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GLYCOSYLTRANSFERASES OF *HELICOBACTER PYLORI* AS A NEW  
TARGET IN PREVENTION AND TREATMENT OF *H. PYLORI* INFECTIONS

5 FIELD OF THE INVENTION

The invention relates to newly identified and isolated polynucleotides and polypeptides of bacterial origin, in particular to novel polynucleotides and polypeptides related to glycosyltransferases involved in biosynthesis of 10 lipopolysaccharides of *Helicobacter pylori*.

BACKGROUND OF THE INVENTION

15 *Helicobacter pylori* is a spiral, microaerophilic, Gram-negative bacterium infecting about 50% of the global human population, and is now recognised as the most common bacterial pathogen of humans worldwide. It is the causative agent of chronic active gastritis in all who harbour it, is responsible for the development of most gastro-duodenal ulcers, and is formally recognised as the carcinogen for 20 certain gastric cancers (Blaser, *Gastroenterology* 102: 720-727 (1992); Parsonnet et al, *N. Engl. J. Med.* 325: 1127-1131 (1991)). *H. pylori* is a highly motile organism and migrates through the superficial mucus layer of the gastric lumen to colonize the underlying gastric pits and associated epithelium. The precise mechanisms by which *H. pylori* injures the gastric mucosa to elicit the 25 aforementioned pathogenic states remains unknown, but it is clear that urease production (Eaton et al, *Infect. Immun.* 59: 2470-2475 (1991)) and motility are required for gastric colonisation of experimental animals. However, the development of gastro-duodenal disease clearly requires additional bacterial virulence factors (Phadnis et al, *Infect. Immun.* 62:1557-1565 (1994); Tummuru et al, *Mol. Microbiol.* 18: 867-876 (1995)). Although several bacterial adhesins 30 and putative receptors on host epithelium have been described (Evans et al, *J. Bacteriol.* 175: 674-683 (1993); Boren et al, *Science* 262: 1892-1895 (1993);

Odenbreit *et al*, *Gut* 37 (Suppl. 1): A1 (1995)), their role in gastric colonization by *H. pylori* has not been clearly established.

Gram-negative bacteria, such as *H. pylori*, have their bacterial cell wall covered with an outer membranous layer consisting of lipids, proteins and lipopolysaccharides (LPS). LPS contain lipid A, a disaccharide of two phosphorylated glucosamine (GlcN) residues with attached fatty acids, and a polysaccharide attached to one of the glucosamine residues through a glycosidic bond. The polysaccharide is composed of a core of approximately 10 sugar residues followed by a repeating series of units of 3 to 5 sugars called the O side chain (O-chain). The number of repeating units in the O-chain varies from about 10 to 40. The sugars found in the O-chain vary among bacterial species, whereas the composition of the core polysaccharide is relatively constant. Lipopolysaccharides are released from bacteria undergoing lysis and are toxic to animals and humans. They are often referred to as endotoxins.

While much attention has focused on the role of bacterial and host proteins in *H. pylori* infection and immunity, the role of LPS in these processes has received less consideration (Moran, *Aliment. Pharmacol. Ther.* 10 (suppl): 39-50 (1996); Yokota *et al*, *Infect. Immun.* 66: 3006-3011 (1998); Wang *et al*, *Mol. Microbiol.* 31: 1265-1274 (1999)). As a major cell surface component, this molecule is well situated to selectively interact with surface components of the host. In particular, LPS could facilitate initial gastric colonisation, be responsible for biological interactions which modify the inflammatory response, and promote a chronic infection.

Comprehensive, detailed structural analysis of *H. pylori* LPS has revealed some unique features of the molecule which may account for certain aspects of *H. pylori*-induced pathogenesis (Aspinall *et al*, *Biochemistry* 35: 2489-2497; 2498-2504 (1996); Aspinall *et al*, *Eur. J. Biochem.* 248: 592-601 (1997); Monteiro *et al*, *J. Biol. Chem.* 273: 11533-11543 (1998)). In addition, *H. pylori* LPS, unlike typical LPS, has low endotoxic properties. Fresh clinical isolates usually display typical smooth type LPS (S-type). The O-chain polysaccharide structure of *H.*

*pylori* type strain (NCTC11637) LPS is composed of a type 2 *N*-acetyllactosamine (LacNAc) chain of various lengths and this O-chain may be partially  $\alpha$ -L-fucosylated or less commonly  $\alpha$ -D-glucosylated or  $\alpha$ -D-galactosylated and may be terminated at the nonreducing end by Lewis blood group epitopes which mimic human cell surface glycoconjugates and glycolipids. However, it remains to be formally established if the O-chain of *H. pylori* LPS contributes to pathogenesis or generates protective immunity. For instance, the Lewis antigens present on the O-chain polysaccharide might reduce the immunogenicity of this molecule during infection, or might trigger autoimmunity. The ability to produce structurally defined truncated LPS molecules should help elucidate the biological role of LPS in *H. pylori* infection and immunity and possibly open a new approach to the treatment and prevention of *H. pylori* infections.

Known methods of prevention and treatment of *H. pylori* infections are either immunogenic or drug-based. The immunogenic approach is mostly intended to provide an immunogenic protection against the bacterium by vaccinating the individual with a usually bacterium-derived immunogen, to elicit an immune response of the organism to future *H. pylori* infections. Among many others, immunogens (antigens) derived from the LPS of *H. pylori* are known in this group of treatments (see, for example, WO 97/14782 and WO 87/07148).

According to the second approach, *H. pylori* infections are treated with antibacterial drugs or combinations of such drugs, intended to eradicate the bacterial population in the infected individual. In this group of treatments, the currently most common are so called triple therapies, in which patients are administered simultaneously two different antibiotics and an acid secretion inhibiting drug. The efficacy of these therapies varies and is often adversely affected by the developing resistance to broad spectrum antibiotics used for this purpose and by side effects of antibiotic therapies, which frequently result in termination of the therapy before completely healing the infection.

In view of the above-indicated deficiencies of the current antibiotic therapies, attempts are made to develop more specific drugs against *H. pylori*, such as

drugs modulating the activity of enzymes specific to the bacteria (see, for example, US 5,801,013 and US 5,942,409). An ideal anti-helicobacterial drug should be selective, meaning that the drug should inhibit *H. pylori* but not the bacterial population of the microflora of the lower intestine. This means that the  
5 molecular target of the drug should be unique to *H. pylori* and/or should be related to its unique phenotypic characteristics, particularly those facilitating the colonization of bacterium's natural ecological niche (the human stomach). While improving the understanding of *H. pylori* pathogenesis, the present invention provides means for developing new anti-helicobacterial drugs possessing such  
10 desirable characteristics.

## SUMMARY OF THE INVENTION

In one aspect, the present invention provides isolated and/or recombinant nucleic acids which encode certain glycosyltransferases of *Helicobacter* origin. The invention also provides recombinant DNA constructs and vectors containing polynucleotide sequences encoding such glycosyltransferases or portions thereof. These nucleic acids and constructs may be used to produce recombinant glycosyltransferases of *Helicobacter* origin by expressing the polynucleotide  
20 sequences in suitable host cells.

In another aspect, the invention provides isolated polypeptides having the enzymatic activity of glycosyltransferases of *Helicobacter* origin. Such polypeptides are useful, among other things, for the identification of modulators, in particular inhibitors of their enzymatic activity, which inhibitors are potential antimicrobial agents. Using the isolated polypeptides of the present invention, potential inhibitors of these enzymes can be screened for antimicrobial or antibiotic effects, without culturing pathogenic strains of *Helicobacter* bacteria, such as *H. pylori*.

According to one embodiment of the invention, preferred glycosyltransferases of *Helicobacter* origin are glycosyltransferases of *H. pylori* involved in the biosynthesis of the bacterial lipopolysaccharide (LPS), in particular of LPS core or LPS O-chain. Disrupting genes of such glycosyltransferases in several strains of

*H. pylori* resulted in mutants unable to complete the structural assembly of LPS and having as a result a reduced ability to colonize the murine stomach.

According to yet another aspect, the present invention provides novel antigens and vaccines used in immunization against *Helicobacter* bacteria, in particular *H. pylori*. The novel antigens are derived from bacteria having deactivated gene of a glycosyltransferase involved in the biosynthesis of the bacterial lipopolysaccharide, in particular of LPS core or LPS O-chain. Purified or partially purified LPS isolated from such mutants is a preferred antigen.

Other advantages, objects and features of the present invention will be readily apparent to those skilled in the art from the following detailed description of preferred embodiments in conjunction with the accompanying drawings and claims.

#### BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 shows amino acid sequence alignment of glycosyltransferases from *H. pylori*, *H. influenzae*, *H. somnus* and *N. meningitidis*. Multiple sequence alignment was performed using the Clustal Alignment Programme (Higgins *et al*, *Gene* 73: 237-244 (1988)). Designations on the left side refer to the origin of the sequences; HP0826 of genebank AE000594 (Tomb *et al*, *Nature* 388:539-547 (1997)), *Haemophilus influenzae* lex2B, U05670 (Cope *et al*, *Mol. Microbiol.* 5: 1113-1124 (1994)), *Haemophilus somnus* lob1, U94833 (Inzana *et al*, *Infect. Immun.* 65: 4675-4681 (1997)) and *Neisseria meningitidis* lgtB, AAC44085 (Jennings *et al*, *Mol. Microbiol.* 18: 729-740 (1995)). Numbers on the right side indicate amino acid positions. Gaps introduced to maximise the alignment are indicated by dashes. Shadings were obtained using the Genedoc Programme ([www.cris.com/~ketchup/genedoc.shtml](http://www.cris.com/~ketchup/genedoc.shtml)). Black indicates 100% identity, dark grey indicates 80% identity, and light grey indicates 60% identity.

Fig. 2 shows a complete FAB-MS spectrum of the methylated intact LPS of 26695::HP0826kan strain.

Fig. 3 is a schematic showing the chemical structure of LPS from parent strains 26695 and SS1 and isogenic mutants of HP0826, HP0159 and HP0479.

Fig. 4 shows results of CZE-MS/MS analysis (+ion mode) of delipidated LPS from 5 *H. pylori* 26695::0159 mutant. Tandem mass spectrum of precursor ions at m/z 902 (doubly protonated ions). Separation conditions: 10 mM ammonium acetate containing 5% methanol, pH 9.0, +25 kV. For MS/MS experiments, nitrogen as a collision gas, E<sub>lab</sub>: 70 eV (laboratory frame of reference).

10 Fig. 5 shows results of CZE-MS/MS (+ion mode) analysis of delipidated LPS from *H. pylori* 0479 mutants. Tandem mass spectrum of precursor ions at m/z 1612. Separation conditions: 10 mM ammonium acetate containing 5% methanol, pH 9.0, +25 kV. For MS/MS experiments, nitrogen as a collision gas, E<sub>lab</sub>: 60 eV (laboratory frame of reference).

15

#### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

As used herein, the terms "identity" and "similarity" mean the degree of sequence 20 relatedness between two or more polynucleotide or polypeptide sequences as determined by the match between strings of such sequences. "Identity" or "similarity" can be readily quantified by algorithms well known to those skilled in the art, implemented in a number of publicly available computer software packages, for example BLAST software package available from NCBI and other 25 sources. The identity or similarity is usually expressed as a percentage of identity with respect to some reference sequence. For example, in a polynucleotide having a sequence 95% identical to a reference nucleotide sequence, 5% of the nucleotides of the reference sequence have been deleted or substituted with another nucleotide, or 5% of another nucleotides have been inserted into the 30 reference sequence. These substitutions, insertions, and/or deletions may take place anywhere between 5' and 3' terminal positions, either interspersed individually among nucleotides of the reference sequence or in one or more contiguous groups within the reference sequence.

The term "isolated" as used herein means altered by the hand of man with respect to its natural state. For a substance occurring in nature, it means that this substance has been changed or removed from its natural environment, or both.

5 For example, a polynucleotide or a polypeptide naturally present in a living organism is not isolated, but the same polynucleotide or polypeptide separated from its natural matrix and coexisting materials is isolated, as the term is employed herein.

10 The term "polynucleotide" or "nucleic acid" refers to any polyribonucleotide or polydeoxyribonucleotide, which may be unmodified or modified RNA or DNA, whether single- or double-stranded. The term "polypeptide" or "protein" refers to any peptide or protein comprising at least two amino acid residues joined to each other by peptide bonds or modified peptide bonds.

15 The term "variant" as used herein means a polynucleotide or polypeptide that differs from a reference polynucleotide or polypeptide but retains its essential properties. A typical variant of a polynucleotide differs in nucleotide sequence from another, reference polynucleotide. Changes in the nucleotide sequence of 20 the variant may or may not alter the amino acid sequence of a polypeptide encoded by the reference polynucleotide. A typical variant of a polypeptide differs in amino acid sequence from another, reference polypeptide. These difference are usually limited and variants of a polypeptide are closely similar overall and identical in many regions. A variant of a polynucleotide or polypeptide may be naturally occurring, such as an allelic variant, or may be prepared by 25 mutagenesis techniques, by direct synthesis, or by other recombinant methods well known to those skilled in the art.

A "fragment" can be considered as a variant of a polynucleotide or polypeptide, 30 having the same nucleotide or amino acid sequence as part of the reference polynucleotide or peptide. A fragment may be "free-standing" or comprised within a larger polynucleotide or polypeptide, normally as a single continuous region.

Nucleic acids referred to herein as "recombinant" are nucleic acids which have been produced by recombinant DNA methodology, including those nucleic acids that are generated by procedures which rely upon a method of artificial recombination, such as polymerase chain reaction (PCR) and/or cloning into a vector using restriction enzymes.

According to one aspect, the invention provides novel isolated polynucleotides and polypeptides, as described in greater detail below. In particular, the invention provides isolated polynucleotides and polypeptides related to glycosyltransferases involved in the biosynthesis of bacterial lipopolysaccharides of bacteria of the genus *Helicobacter*, more particularly the lipopolysaccharides of the species *Helicobacter pylori* and various strains thereof. In a preferred embodiment, the glucosyltransferases as those involved in the biosynthesis of the bacterial LPS, in particular of LPS core or LPS O-chain. Most particularly, the invention provides isolated polynucleotides and polypeptides identical over their entire lengths to sequences set out in Table 1.

Table 1. Polynucleotide and polypeptide sequences

Sequences from strain 26695 of *H. pylori*

## 5 A. polynucleotide sequence: ORF HP0826 [SEQ ID NO:1]

	ttgcgtgttt ttgccatttc tttaaatcaa aaagtgtgcg atacatttg 60
	agagacacca caacttact caatagcatc aatgccaccc accaccaagc gcaaatttt 120
10	gatgcgattt attctaaaac ttttgaaggc gggttgcacc ccttagtcaa aaagcattta 180
	cacccttatt tcatacgcgca aaacatcaa gacatgggta ttacaaccaa tctcatcagt 240
	gaggtttcta agttttatta cgctttaaaa taccatgcga agtttatgag cttgggggag 300
	cttgggtgct atgcgagtca ttattccttg tggaaaaat gcatagaact caatgaagcg 360
	atctgtattt tagaagacga tataaccttg aaagaggatt ttaaagaggg cttggatttt 420
15	ttagaaaaac acatccaaga gttaggctat atccgcttga tgcatattt gtatgatgcc 480
	agtgtaaaaa gtgagccatt gaggcataaa aaccacgaga tacaagagcg tgtgggatc 540
	attaaagctt atagcgaagg ggtggggact caaggctatg tgatcacgccc taagattgcc 600
	aaagttttt tgaaatgcag ccgaaaatgg gttgttccctg tggatacgat aatggacgct 660
	acttttatcc atggcgtgaa aaatctggtg ttacaacctt ttgtgatcgc tgatgatgag 720
20	caaatctcta cgatagcag aaaagaagaa ccttatagcc cttaaatcgc cttaatgaga 780
	gaactccatt taaaatattt gaaatattgg cagtttgtat aa 822

## B. polypeptide sequence deduced from sequence A [SEQ ID NO:2]

25	Leu Arg Val Phe Ala Ile Ser Leu Asn Gln Lys Val Cys Asp Thr Phe 1 5 10 15
	Gly Leu Val Phe Arg Asp Thr Thr Leu Leu Asn Ser Ile Asn Ala 20 25 30
30	Thr His His Gln Ala Gln Ile Phe Asp Ala Ile Tyr Ser Lys Thr Phe 35 40 45
	Glu Gly Gly Leu His Pro Leu Val Lys Lys His Leu His Pro Tyr Phe 50 55 60
	Ile Thr Gln Asn Ile Lys Asp Met Gly Ile Thr Thr Asn Leu Ile Ser 65 70 75 80
35	Glu Val Ser Lys Phe Tyr Tyr Ala Leu Lys Tyr His Ala Lys Phe Met 85 90 95
	Ser Leu Gly Glu Leu Gly Cys Tyr Ala Ser His Tyr Ser Leu Trp Glu 100 105 110
40	Lys Cys Ile Glu Leu Asn Glu Ala Ile Cys Ile Leu Glu Asp Asp Ile 115 120 125
	Thr Leu Lys Glu Asp Phe Lys Glu Gly Leu Asp Phe Leu Glu Lys His 130 135 140
	Ile Gln Glu Leu Gly Tyr Ile Arg Leu Met His Leu Leu Tyr Asp Ala 145 150 155 160
45	Ser Val Lys Ser Glu Pro Leu Ser His Lys Asn His Glu Ile Gln Glu 165 170 175
	Arg Val Gly Ile Ile Lys Ala Tyr Ser Glu Gly Val Gly Thr Gln Gly 180 185 190
50	Tyr Val Ile Thr Pro Lys Ile Ala Lys Val Phe Leu Lys Cys Ser Arg 195 200 205
	Lys Trp Val Val Pro Val Asp Thr Ile Met Asp Ala Thr Phe Ile His 210 215 220
	Gly Val Lys Asn Leu Val Leu Gln Pro Phe Val Ile Ala Asp Asp Glu 225 230 235 240
55	Gln Ile Ser Thr Ile Ala Arg Lys Glu Glu Pro Tyr Ser Pro Lys Ile 245 250 255

Ala Leu Met Arg Glu Leu His Phe Lys Tyr Leu Lys Tyr Trp Gln Phe  
                  260                 265                 270  
 Val

5

C. polynucleotide sequence: ORF HP0159 [SEQ ID NO:3]

	atgaggattta ttattccttat tgtcatcgct tttgataatc actatgccat gccggctggc 60
10	gtgagcttgc attccatgct agcttgcgt aaaacagaac acccccatac acaaaaatgat 120
	agtggaaaac ttttttataa gatccactgc ctgggtggata acttaagcct tgaaaaccag 180
	agcaaaactaa aagagactct agcccccttt agcgcttttt cgagcctaga attttttagac 240
	attcaaccc ccaatcttca cgccactcca atagaaccct ctgcgattga taaaatcaat 300
15	gaagctttt tgcaactcaa tatttacgct aagactcgct tttctaaaat ggtcatgtgc 360
	cgcttgttt tggcttcctt attcccacaa tacgacaaaa tcatcatgtt tgatgcagac 420
	actttgttt taaacgatgt gagcgagagc ttttcattcc cactagatgg ctattatttt 480
	ggagcggcta aagatttgc ttccgataaaa agccctaaac attttcaaata agtgcgagaaa 540
	aaagaccctc gtcaagcctt ttccctttat gagcattacc ttaatgaaag cgatatgcaa 600
20	atcatctatg aaagcaatta taacgcccggg ttttagtcg tgaatttaaa gctgtggcgt 660
	gctgatcat ttagaagagcg cttactcaat ttaacccatc aaaaaggcca gtgcgtgtt 720
	taccctgaac aggaccttt aacgctcgca tgctatcaaa aagttttaat cttgccttat 780
	atttataaca cccacccctt catggccaat caaaaacgct tcatccctga caaaaaagaaa 840
	atcgctcatgc tgcatttta tttttagga aaaccttggg ttttacctac ttttcatat 900
25	tctaaagaat ggcatgagac tcttttaaaa acccctttt atgctgaata ttccgtgaaa 960
	ttccttaaac aaatgacaga atgtttaagc cttaaagaca aaaaaaaaaac ctttgaattt 1020
	cttgcccccc tactcaataa aaaaaccctt tttagaatacg tcttttttag attgaatagg 1080
	attttcaaac qcttaaaaaga aaaattttt aactcttag 1119

D. polypeptide sequence deduced from sequence C [SEQ ID NO:4]

30

	Met	Ser	Ile	Ile	Ile	Pro	Ile	Val	Ile	Ala	Phe	Asp	Asn	His	Tyr	Ala
1							5				10					15
	Met	Pro	Ala	Gly	Val	Ser	Leu	Tyr	Ser	Met	Leu	Ala	Cys	Ala	Lys	Thr
							20			25					30	
35	Glu	His	Pro	Gln	Ser	Gln	Asn	Asp	Ser	Glu	Lys	Leu	Phe	Tyr	Lys	Ile
							35		40					45		
	His	Cys	Leu	Val	Asp	Asn	Leu	Ser	Leu	Glu	Asn	Gln	Ser	Lys	Leu	Lys
							50		55			60				
40	Glu	Thr	Leu	Ala	Pro	Phe	Ser	Ala	Phe	Ser	Ser	Leu	Glu	Phe	Leu	Asp
						65		70			75				80	
	Ile	Ser	Thr	Pro	Asn	Leu	His	Ala	Thr	Pro	Ile	Glu	Pro	Ser	Ala	Ile
							85			90					95	
	Asp	Lys	Ile	Asn	Glu	Ala	Phe	Leu	Gln	Leu	Asn	Ile	Tyr	Ala	Lys	Thr
						100			105			110				
45	Arg	Phe	Ser	Lys	Met	Val	Met	Cys	Arg	Leu	Phe	Leu	Ala	Ser	Leu	Phe
						115			120			125				
	Pro	Gln	Tyr	Asp	Lys	Ile	Ile	Met	Phe	Asp	Ala	Asp	Thr	Leu	Phe	Leu
						130			135			140				
50	Asn	Asp	Val	Ser	Glu	Ser	Phe	Phe	Ile	Pro	Leu	Asp	Gly	Tyr	Tyr	Phe
							145		150			155			160	
	Gly	Ala	Ala	Lys	Asp	Phe	Ala	Ser	Asp	Lys	Ser	Pro	Lys	His	Phe	Gln
							165			170			175			
	Ile	Val	Arg	Glu	Lys	Asp	Pro	Arg	Gln	Ala	Phe	Ser	Leu	Tyr	Glu	His
							180			185			190			
55	Tyr	Leu	Asn	Glu	Ser	Asp	Met	Gln	Ile	Ile	Tyr	Glu	Ser	Asn	Tyr	Asn
							195		200			205				
	Ala	Gly	Phe	Leu	Val	Val	Asn	Leu	Lys	Leu	Trp	Arg	Ala	Asp	His	Leu
							210		215			220				

Glu Glu Arg Leu Leu Asn Leu Thr His Gln Lys Gly Gln Cys Val Phe  
 225 230 235 240  
 Tyr Pro Glu Gln Asp Leu Leu Thr Leu Ala Cys Tyr Gln Lys Val Leu  
 245 250 255  
 5 Ile Leu Pro Tyr Ile Tyr Asn Thr His Pro Phe Met Ala Asn Gln Lys  
 260 265 270  
 Arg Phe Ile Pro Asp Lys Lys Glu Ile Val Met Leu His Phe Tyr Phe  
 275 280 285  
 Val Gly Lys Pro Trp Val Leu Pro Thr Phe Ser Tyr Ser Lys Glu Trp  
 10 290 295 300  
 His Glu Thr Leu Leu Lys Thr Pro Phe Tyr Ala Glu Tyr Ser Val Lys  
 305 310 315 320  
 Phe Leu Lys Gln Met Thr Glu Cys Leu Ser Leu Lys Asp Lys Gln Lys  
 325 330 335  
 15 Thr Phe Glu Phe Leu Ala Pro Leu Leu Asn Lys Lys Thr Leu Leu Glu  
 340 345 350  
 Tyr Val Phe Phe Arg Leu Asn Arg Ile Phe Lys Arg Leu Lys Glu Lys  
 355 360 365  
 20 Phe Phe Asn Ser  
 370

E. polynucleotide sequence: ORF HP0479 [SEQ ID NO:5]

25 atgcatgttg cttgtctttt ggcttttaggg gataatctca tcacgcttag ccttttaaaa 60  
 gaaatcgctt tcaaacagca acaacccctt aaaatcctag gtactcgctt gactttaaaa 120  
 atcgccaagc ttttagaatg cgaaaaaacat tttgaaatca ttctctttt tgaaaatgtc 180  
 cctgctttttt atgaccttaa aaaacaaggc gtttttttgg cgatgaagga ttttttatgg 240  
 ttgttaaaag cgattaaaaaa gcataaatac aaacgtttga ttttggaaaa acaggatttt 300  
 30 agaagcactt ttttagccaa attcattccc ataaccactc caaataaaaga aattaaaaaac 360  
 gtttatcaaa accgccagga gttgtttctt caaattttagt ggcatgttt tgataacccc 420  
 ccatatcccc tgaattttaaa aaaccccaaa aagattttga tcaacccttt cacaagatcc 480  
 atagaccgaa gtatcccttt agagcattta caaatcgttt taaaactttt aaaacccttt 540  
 tttgttacgc ttttagattt tgaagaacga tacgctttt taaaagacag agtcgctcat 600  
 35 tatcgcgcta aaaccagttt agaagaagtt aaaaacctga ttttagaaag cgatttgtat 660  
 ataggagggg attcgttttt gatccatttg gcttactatt taaaagaaaaa ttattttatc 720  
 ttttttata gggataatga tgatttcatg ccgcctaata gtaagaataa aaattttcta 780  
 aaagcccaca aaagccattc tatagaacaa gatttagcca aaaaattccg ccatttgggg 840  
 ctattataa 849

F. polypeptide sequence deduced from sequence E [SEQ ID NO:6]

40 Met His Val Ala Cys Leu Leu Ala Leu Gly Asp Asn Leu Ile Thr Leu  
 1 5 10 15  
 Ser Leu Leu Lys Glu Ile Ala Phe Lys Gln Gln Pro Leu Lys Ile  
 20 25 30  
 Leu Gly Thr Arg Leu Thr Leu Lys Ile Ala Lys Leu Leu Glu Cys Glu  
 35 40 45  
 45 Lys His Phe Glu Ile Ile Pro Leu Phe Glu Asn Val Pro Ala Phe Tyr  
 50 55 60  
 Asp Leu Lys Lys Gln Gly Val Phe Leu Ala Met Lys Asp Phe Leu Trp  
 65 70 75 80  
 Leu Leu Lys Ala Ile Lys Lys His Gln Ile Lys Arg Leu Ile Leu Glu  
 55 85 90 95  
 Lys Gln Asp Phe Arg Ser Thr Phe Leu Ala Lys Phe Ile Pro Ile Thr  
 100 105 110  
 Thr Pro Asn Lys Glu Ile Lys Asn Val Tyr Gln Asn Arg Gln Glu Leu

	115		120		125											
	Phe	Ser	Gln	Ile	Tyr	Gly	His	Val	Phe	Asp	Asn	Pro	Pro	Tyr	Pro	Met
	130						135					140				
5	Asn	Leu	Lys	Asn	Pro	Lys	Lys	Ile	Leu	Ile	Asn	Pro	Phe	Thr	Arg	Ser
	145					150					155			160		
	Ile	Asp	Arg	Ser	Ile	Pro	Leu	Glu	His	Leu	Gln	Ile	Val	Leu	Lys	Leu
							165			170				175		
	Leu	Lys	Pro	Phe	Cys	Val	Thr	Leu	Leu	Asp	Phe	Glu	Glu	Arg	Tyr	Ala
10						180				185				190		
	Phe	Leu	Lys	Asp	Arg	Val	Ala	His	Tyr	Arg	Ala	Lys	Thr	Ser	Leu	Glu
						195			200				205			
	Glu	Val	Lys	Asn	Leu	Ile	Leu	Glu	Ser	Asp	Leu	Tyr	Ile	Gly	Gly	Asp
						210			215				220			
15	Ser	Phe	Leu	Ile	His	Leu	Ala	Tyr	Tyr	Leu	Lys	Lys	Asn	Tyr	Phe	Ile
						225			230			235			240	
	Phe	Phe	Tyr	Arg	Asp	Asn	Asp	Asp	Phe	Met	Pro	Pro	Asn	Ser	Lys	Asn
							245			250				255		
	Lys	Asn	Phe	Leu	Lys	Ala	His	Lys	Ser	His	Ser	Ile	Glu	Gln	Asp	Leu
20							260			265				270		
	Ala	Lys	Lys	Phe	Arg	His	Leu	Gly	Leu	Leu						
						275			280							

## G. polynucleotide sequence: ORF 1191

[SEQ ID NO:7]

H<sub>n</sub> polypeptide sequence deduced from sequence C [SEQ ID NO:8]

	Met	Ser	Val	Asn	Ala	Pro	Lys	Arg	Met	Arg	Ile	Leu	Leu	Arg	Leu	Pro
	1					5				10					15	
50	Asn	Trp	Leu	Gly	Asp	Gly	Val	Met	Ala	Ser	Ser	Leu	Phe	Tyr	Thr	Leu
						20				25				30		
	Lys	His	His	Tyr	Pro	Asn	Ala	His	Phe	Ile	Leu	Val	Gly	Pro	Thr	Ile
						35			40				45			
55	Thr	Cys	Glu	Leu	Phe	Lys	Lys	Asp	Glu	Lys	Ile	Glu	Ala	Val	Phe	Ile
						50			55			60				
	Asp	Asn	Thr	Lys	Lys	Ser	Phe	Phe	Arg	Leu	Leu	Ala	Ile	His	Lys	Leu
						65			70			75			80	
	Ala	Gln	Lys	Ile	Gly	Arg	Cys	Asp	Ile	Ala	Ile	Thr	Leu	Asn	Asn	His
						85			90					95		

Phe Tyr Ser Ala Phe Leu Leu Tyr Ala Thr Lys Thr Pro Val Arg Ile  
                  100                 105                 110  
 Gly Phe Ala Gln Phe Phe Arg Ser Leu Phe Leu Ser His Ala Ile Ala  
                  115                 120                 125  
 5 Pro Ala Pro Lys Glu Tyr His Gln Val Glu Lys Tyr Cys Phe Leu Phe  
                  130                 135                 140  
 Ser Gln Phe Leu Glu Lys Glu Leu Asp Gln Lys Ser Val Leu Pro Leu  
                  145                 150                 155                 160  
 Lys Leu Ala Phe Asn Leu Pro Thr His Thr Pro Asn Thr Pro Lys Lys  
 10                 165                 170                 175  
 Ile Gly Phe Asn Pro Ser Ala Ser Tyr Gly Ser Ala Lys Arg Trp Pro  
                  180                 185                 190  
 Ala Ser Tyr Tyr Ala Glu Val Ser Ala Val Leu Leu Glu Lys Gly His  
                  195                 200                 205  
 15 Glu Ile Tyr Phe Phe Gly Ala Lys Glu Asp Ala Ile Val Ser Glu Glu  
                  210                 215                 220  
 Ile Leu Lys Leu Ile Lys Gly Ser Leu Lys Asn Pro Ser Leu Phe His  
                  225                 230                 235                 240  
 Asn Ala Tyr Asn Leu Cys Gly Lys Thr Ser Ile Glu Glu Leu Ile Glu  
 20                 245                 250                 255  
 Arg Ile Ala Val Leu Asp Leu Phe Ile Thr Asn Asp Ser Gly Pro Met  
                  260                 265                 270  
 His Val Ala Ala Ser Met Gln Thr Pro Leu Ile Ala Leu Phe Gly Pro  
                  275                 280                 285  
 25 Thr Asp Glu Lys Glu Thr Arg Pro Tyr Lys Ala Gln Lys Thr Ile Val  
                  290                 295                 300  
 Leu Asn His His Leu Ser Cys Ala Pro Cys Lys Lys Arg Val Cys Pro  
                  305                 310                 315                 320  
 Leu Lys Asn Ala Lys Asn His Leu Cys Met Lys Ser Ile Thr Pro Leu  
 30                 325                 330                 335  
 Glu Val Leu Glu Ala Ala His Thr Leu Leu Glu Glu Pro  
                  340                 345

35 Sequences from strain SS1 of *H. pylori*

## I. polynucleotide sequence: ORF SS0826 [SEQ ID NO:9]

40 ttgcgtat ttatcatttc tttaaatcaa aaagtgtgcg ataaaattgg tttgggtttt 60  
 agagacacca cgactttact caatagcatc aatgccaccc accaccaagt gcaaattttt 120  
 gatgcgattt attctaaaac ttttgaaggc gggttgcacc ccttagtgaa aaagcattta 180  
 cacccttatt tcatacgc aAACATCAAAC gacatggaa ttacaaccag tctcatcagt 240  
 gaggtttcta agttttatta cgctttaaaa taccatgcga agtttatgag cttgggagag 300  
 cttgggtgct atgcgagcca ttattccttg tggaaaaat gcatagaact caatgaagcg 360  
 45 atctgtat tt tagaagacga tataaccttgg aaagaggatt taaaagaggg cttggat 420  
 ttagaaaaac acatccaaga gttaggctat gttcgcttga tgcattttt atatgatccc 480  
 aatattaaaa gtgagccatt gaaccataaa aaccacgaga tacaagagcg tgtaggatt 540  
 attaaagctt atagcgaagg ggtggggact caaggctatg tgatcacgccc caagattgcc 600  
 aaagttttta aaaaacacag ccgaaaaatgg gttgttcctg tggatacgat aatggacgct 660  
 50 acttttatcc atggcgtgaa aaatctggtg ttacaacctt ttgtgatcgc tgatgatgag 720  
 caaatctcta cgatagcgcg aaaagaacaa ccttatacgcc cttaaatcgc cttaatgaga 780  
 gaactccatt ttaaatattt gaaatattgg cagtttatat ag 822

## 55 J. polypeptide sequence deduced from sequence I [SEQ ID NO:10]

Leu Arg Ile Phe Ile Ile Ser Leu Asn Gln Lys Val Cys Asp Lys Phe  
                  1                 5                 10                 15

35 K. polynucleotide sequence: ORF SS0159 [SEQ ID NO:11]

	atgaggattta	ctattccttat	tgttatcgct	tttacaacatc	attacgccat	tccggctggc	60
	gtgagccctgt	attccatgct	agcttgcaact	aaaacagaac	accccccaatc	acaaaatgat	120
40	agtaaaaaac	ttttttataaa	aatccactgc	ctggtagata	acttaaagcct	tgaaaaccag	180
	tgcaaattga	aagaaactct	agcccccttt	agcgctttta	tgagcgtgga	tttttttagac	240
	atttcaaccc	ctaattttta	caccccttca	atagaaccct	ctgcgattga	taaaaatcaat	300
	gaagctttt	tgcaactcaa	tatttacgct	aagactcgct	tttctaaaat	ggtcatgtgc	360
45	cgcttggttt	tggcctcttt	attcccgc当地	tacgacaaaa	tcatcatgtt	tgatgc当地	420
	actttgtttt	taaacgatgt	gagcgagagt	ttttttatcc	cgctagatgg	ttattat	480
	ggagcggctta	aagattttc	ttctcctaaa	aaccttaaac	atttcaaac	agaaaaggag	540
	agagagccctc	gccaaaaatt	tttctccat	gaggcattacc	ttaaagaaaa	agacatgaaa	600
	atcatttgtg	aaaaccacta	taatgttggg	ttttaatcg	tgaattaaa	gctgtggcgt	660
50	gctgatcatt	tagaagaacg	cttactcaat	ttaacccatc	aaaaaggcca	gtgtgtgtt	720
	tgccctgaac	aggatatttt	aacgctcgca	tgc当地atcaa	aagtttaca	attacctt	780
	atttacaaca	cccacccctt	catggtcaat	caaaaacgct	tcatccctaa	caaaaaagaa	840
	atcgatgc	tgc当地tttta	ttttaggaa	aaaccttggg	tttacccac	tgcttata	900
	tctaaagaat	ggcatgagac	tctttaaaa	accctttt	acgctgaata	ttccgtgaaa	960
55	tttcttaaac	aaatgacaga	attttaagc	cttaaagaca	aacaaaaaaac	cttgaattt	1020
	cttgc当地cc	tactcaataa	aaaaaccctt	ttagaatatg	tcttttttag	attgaatagg	1080
	attttcaaaac	gctaaaaga	aaaactttt	aactcttagc			1120

## L. polypeptide sequence deduced from sequence K [SEQ ID NO:12]

Met Ser Ile Thr Ile Pro Ile Val Ile Ala Phe Asp Asn His Tyr Ala  
 1 5 10 15  
 Ile Pro Ala Gly Val Ser Leu Tyr Ser Met Leu Ala Cys Thr Lys Thr  
 20 25 30  
 Glu His Pro Gln Ser Gln Asn Asp Ser Glu Lys Leu Phe Tyr Lys Ile  
 35 40 45  
 His Cys Leu Val Asp Asn Leu Ser Leu Glu Asn Gln Cys Lys Leu Lys  
 50 55 60  
 Glu Thr Leu Ala Pro Phe Ser Ala Phe Met Ser Val Asp Phe Leu Asp  
 65 70 75 80  
 Ile Ser Thr Pro Asn Leu Tyr Thr Pro Ser Ile Glu Pro Ser Ala Ile  
 85 90 95  
 15 Asp Lys Ile Asn Glu Ala Phe Leu Gln Leu Asn Ile Tyr Ala Lys Thr  
 100 105 110  
 Arg Phe Ser Lys Met Val Met Cys Arg Leu Phe Leu Ala Ser Leu Phe  
 115 120 125  
 Pro Gln Tyr Asp Lys Ile Ile Met Phe Asp Ala Asp Thr Leu Phe Leu  
 130 135 140  
 Asn Asp Val Ser Glu Ser Phe Phe Ile Pro Leu Asp Gly Tyr Tyr Phe  
 145 150 155 160  
 Gly Ala Ala Lys Asp Phe Ser Ser Pro Lys Asn Leu Lys His Phe Gln  
 165 170 175  
 25 Thr Glu Arg Glu Arg Glu Pro Arg Gln Lys Phe Phe Leu His Glu His  
 180 185 190  
 Tyr Leu Lys Glu Lys Asp Met Lys Ile Ile Cys Glu Asn His Tyr Asn  
 195 200 205  
 Val Gly Phe Leu Ile Val Asn Leu Lys Leu Trp Arg Ala Asp His Leu  
 210 215 220  
 Glu Glu Arg Leu Leu Asn Leu Thr His Gln Lys Gly Gln Cys Val Phe  
 225 230 235 240  
 Cys Pro Glu Gln Asp Ile Leu Thr Leu Ala Cys Tyr Gln Lys Val Leu  
 245 250 255  
 35 Gln Leu Pro Tyr Ile Tyr Asn Thr His Pro Phe Met Val Asn Gln Lys  
 260 265 270  
 Arg Phe Ile Pro Asn Lys Lys Glu Ile Val Met Leu His Phe Tyr Phe  
 275 280 285  
 Val Gly Lys Pro Trp Val Leu Pro Thr Ala Leu Tyr Ser Lys Glu Trp  
 290 295 300  
 His Glu Thr Leu Leu Lys Thr Pro Phe Tyr Ala Glu Tyr Ser Val Lys  
 305 310 315 320  
 Phe Leu Lys Gln Met Thr Glu Phe Leu Ser Leu Lys Asp Lys Gln Lys  
 325 330 335  
 45 Thr Phe Glu Phe Leu Ala Pro Leu Leu Asn Lys Lys Thr Leu Leu Glu  
 340 345 350  
 Tyr Val Phe Phe Arg Leu Asn Arg Ile Phe Lys Arg Leu Lys Glu Lys  
 355 360 365  
 Leu Leu Asn Ser  
 370

## M. polynucleotide sequence: ORF SS0479 [SEQ ID NO:13]

55 atgcatgttg cttgtctttt ggcttttaggg gataaacctca tcacgcttag cctttgtgaa 60  
 gaaatcgctc tcaaacagca acaacccctt aaaatcctag gtactcgttt gactttaaaa 120  
 atcgccaaggc ttttagaatg cgaaaaaacat tttgaaatca ttccctgtttt taaaaatatc 180  
 cccgctttttt atgaccttaa aaaacaaggc gttttttggg cgatgaagga ttttttatgg 240

ttattaaaag cgcttaagaa gcacaaaatc aaacacttga ttttagaaaa acaagattt 300  
 agaagcgctc ttttatccaa atttgttcc ataaccactc caaataaaga aattaaaaat 360  
 gcttattcaaa accgccagga gttgtttct caaattttatg ggcatgttt tgataatagt 420  
 caatattcca tgagttaaa aaaccccaa aagattttaa tcaaccctt cacgagagaa 480  
 5 aataatagaa atatttctt agaacatttga caaatcggtt taaaactgtt aaaaccctt 540  
 tgtgttacgc ttttagattt tgaagaacga tacgctttt taaaagatga agtcgctcat 600  
 tatcgcgcta aaaccagttt agaagaagct aaaaacctga ttttagaaag cgatttgat 660  
 ataggggggg attcggtttt gatccatttga gtttactatt taaagaaaaa ttattttac 720  
 ttttttata gggataatga cgatttcattt ccgcctaaga atgaaaattt tctaaaagcc 780  
 10 cataaaagcc atttcataga gcaggatttga gccacccagt tccgcccattt gggctattt 840  
 taa 843

### N. polypeptide sequence deduced from sequence M [SEQ ID NO:14]

15 Met His Val Ala Cys Leu Leu Ala Leu Gly Asp Asn Leu Ile Thr Leu  
 1 5 10 15  
 Ser Leu Cys Glu Glu Ile Ala Leu Lys Gln Gln Pro Leu Lys Ile  
 20 25 30  
 20 Leu Gly Thr Arg Leu Thr Leu Lys Ile Ala Lys Leu Leu Glu Cys Glu  
 35 40 45  
 Lys His Phe Glu Ile Ile Pro Val Phe Lys Asn Ile Pro Ala Phe Tyr  
 50 55 60  
 Asp Leu Lys Lys Gln Gly Val Phe Trp Ala Met Lys Asp Phe Leu Trp  
 25 65 70 75 80  
 Leu Leu Lys Ala Leu Lys Lys His Lys Ile Lys His Leu Ile Leu Glu  
 85 90 95  
 Lys Gln Asp Phe Arg Ser Ala Leu Leu Ser Lys Phe Val Ser Ile Thr  
 100 105 110  
 30 Thr Pro Asn Lys Glu Ile Lys Asn Ala Tyr Gln Asn Arg Gln Glu Leu  
 115 120 125  
 Phe Ser Gln Ile Tyr Gly His Val Phe Asp Asn Ser Gln Tyr Ser Met  
 130 135 140  
 Ser Leu Lys Asn Pro Lys Lys Ile Leu Ile Asn Pro Phe Thr Arg Glu  
 35 145 150 155 160  
 Asn Asn Arg Asn Ile Ser Leu Glu His Leu Gln Ile Val Leu Lys Leu  
 165 170 175  
 Leu Lys Pro Phe Cys Val Thr Leu Leu Asp Phe Glu Glu Arg Tyr Ala  
 180 185 190  
 40 Phe Leu Lys Asp Glu Val Ala His Tyr Arg Ala Lys Thr Ser Leu Glu  
 195 200 205  
 Glu Ala Lys Asn Leu Ile Leu Glu Ser Asp Leu Tyr Ile Gly Gly Asp  
 210 215 220  
 Ser Phe Leu Ile His Leu Ala Tyr Tyr Leu Lys Lys Asn Tyr Phe Ile  
 45 225 230 235 240  
 Phe Phe Tyr Arg Asp Asn Asp Asp Phe Met Pro Pro Lys Asn Glu Asn  
 245 250 255  
 Phe Leu Lys Ala His Lys Ser His Phe Ile Glu Gln Asp Leu Ala Thr  
 260 265 270  
 50 Gln Phe Arg His Leu Gly Leu Leu  
 275 280

### Sequences from strain PJ1 of *H. pylori*

55

### O. polynucleotide sequence: ORF PJ1-0479 [SEQ ID NO:15]

atgcatgttg cttgttttt ggctttaggg gataacctca tcacgcttag cctttaaaaa 60  
 gaaatcgctt ccaaacagca acggccccctt aaaatccttag gcactcggtt gactttaaaaa 120

atcgccaagc ttttagaatg cgaaaaacat tttgaaatca ttccctatgg tgaaaatatac 180  
 cctgcctttt atgatcttaa aaaacaaggc gtttttggg cgatgaagga ttttttatgg 240  
 ttgttaaaag caattaagaa gcacaaaatc aaacatttga ttttagaaaa acaagattt 300  
 agaagtttc ttttatccaa atttgttcc ataaccactc ccaataaaaga aattaaaaac 360  
 5 gtttatcaaa accgccagga gttgtttct ccaatttatg ggcatgttt tgataacccc 420  
 ccataccca tgaatttaaa aaaccccaa aagatttga tcaaccctt cacaagatcc 480  
 atagagcgaa gtatccctt agagcattt aaaaatcggtt taaaactctt aaaacccttt 540  
 tgtgttacgc ttttagattt tgaagaacga tacgctttt tacaaaatga agccactcat 600  
 tatacggtcta aaaccagttt agaagaagtt aaaagcctga ttttagaaag cgatttgat 660  
 10 ataggggggg attcgtttt aatccatgg gcttactatt taaagaaaaa ttatttatc 720  
 ttttttata ggataatga cgatttcatg ccacctaatg gtaagaagga aaatttcta 780  
 aaagccaca aaagccatta catagaacag gatttagcca aaaaattccg ccatttgggg 840  
 cttattataa 850

15

P. polypeptide sequence deduced from sequence O [SEQ ID NO:16]

	Met His Val Ala Cys Leu Leu Ala Leu Gly Asp Asn Leu Ile Thr Leu
20	1 5 10 15
	Ser Leu Leu Lys Glu Ile Ala Ser Lys Gln Gln Arg Pro Leu Lys Ile
	20 25 30
	Leu Gly Thr Arg Leu Thr Leu Lys Ile Ala Lys Leu Leu Glu Cys Glu
	35 40 45
25	Lys His Phe Glu Ile Ile Pro Ile Phe Glu Asn Ile Pro Ala Phe Tyr
	50 55 60
	Asp Leu Lys Lys Gln Gly Val Phe Trp Ala Met Lys Asp Phe Leu Trp
	65 70 75 80
30	Leu Leu Lys Ala Ile Lys Lys His Lys Ile Lys His Leu Ile Leu Glu
	85 90 95
	Lys Gln Asp Phe Arg Ser Phe Leu Ser Lys Phe Val Ser Ile Thr
	100 105 110
	Thr Pro Asn Lys Glu Ile Lys Asn Val Tyr Gln Asn Arg Gln Glu Leu
	115 120 125
35	Phe Ser Pro Ile Tyr Gly His Val Phe Asp Asn Pro Pro Tyr Pro Met
	130 135 140
	Asn Leu Lys Asn Pro Lys Lys Ile Leu Ile Asn Pro Phe Thr Arg Ser
	145 150 155 160
40	Ile Glu Arg Ser Ile Pro Leu Glu His Leu Lys Ile Val Leu Lys Leu
	165 170 175
	Leu Lys Pro Phe Cys Val Thr Leu Leu Asp Phe Glu Glu Arg Tyr Ala
	180 185 190
	Phe Leu Gln Asn Glu Ala Thr His Tyr Arg Ala Lys Thr Ser Leu Glu
	195 200 205
45	Glu Val Lys Ser Leu Ile Leu Glu Ser Asp Leu Tyr Ile Gly Gly Asp
	210 215 220
	Ser Phe Leu Ile His Leu Ala Tyr Tyr Leu Lys Lys Asn Tyr Phe Ile
	225 230 235 240
	Phe Phe Tyr Arg Asp Asn Asp Asp Phe Met Pro Pro Asn Gly Lys Lys
50	245 250 255
	Glu Asn Phe Leu Lys Ala His Lys Ser His Tyr Ile Glu Gln Asp Leu
	260 265 270
	Ala Lys Lys Phe Arg His Leu Gly Leu Ile Ile
	275 280

55

Preferred embodiments of the invention are polynucleotides coding for *H. pylori* glycosyltransferases involved in the biosynthesis of the core or O-chain regions of the bacterial lipopolysaccharide (LPS), in particular polynucleotides having sequences shown in Table 1 (SEQ ID NO: 1, 3, 5, 7, 9, 11, 13 and 15),  
5 polynucleotides closely related thereto, as well as fragments and variants thereof.  
Another preferred embodiments of the invention are polynucleotides that are at  
least 70% identical over their entire length to polynucleotides shown in Table 1,  
preferably at least 80% identical, more preferably at least 90% identical, most  
preferably at least 95% identical, and polynucleotides that are complementary to  
10 such polynucleotides. Furthermore, those with at least 97% are highly preferred  
among those with at least 95%, and among these those with at least 98% and at  
least 99% are particularly highly preferred, with at least 99% being the most  
preferred.

15 Of the polynucleotides showing substantial identity to the polynucleotides shown  
in Table 1 (SEQ ID NO: 1, 3, 5, 7, 9, 11, 13 and 15), preferred are those which  
encode polypeptides showing substantially the same biological function or activity  
as the polypeptides shown in Table 1 (SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14 and  
16).

20 Polynucleotides shown in Table 1 correspond to open reading frames HP0826  
(SEQ ID NO: 1), HP0159 (SEQ ID NO: 3), HP0479 (SEQ ID NO: 5) and HP1191  
(SEQ ID NO: 7) of the genomic DNA of *H. pylori* strain 26695, to open reading  
frames SS0826 (SEQ ID NO: 9), SS0159 (SEQ ID NO: 11) and SS0479 (SEQ ID  
NO: 13) of the genomic DNA of *H. pylori* strain SS1, and to open reading frame  
25 PJ1-0479 (SEQ ID NO: 15) of the genomic DNA of *H. pylori* strain PJ1. Among  
several others, ORFs HP0826, HP0159, HP0479 and HP1191 have been  
identified using the complete annotated genome sequence of *H. pylori* strain  
26695 and BLAST analysis as potentially coding for glycosyltransferases. They  
30 have been proven, directly or indirectly, to encode a  $\beta$ -1,4-galactosyltransferase  
(HP0826), a  $\alpha$ -1,6-glucosyltransferase (HP0159), a heptosyltransferase  
(HP0479), and an ADP-heptose-LPS heptosyltransferase II (HP1191), which are  
enzymes involved in the biosynthesis of the *H. pylori* lipopolysaccharide. ORFs

identified by BLAST analysis have been cloned, expressed, and isolated using techniques well known to those skilled in the art, also discussed more in detail further in this disclosure.

5     The isolated polynucleotides of the present invention can be used in the production of polypeptides they encode. For example, a polynucleotide containing all or part of the coding sequence for a *Helicobacter* glycosyltransferase can be incorporated into various DNA constructs, such as expression cassettes, and vectors, such as recombinant plasmids, adapted for  
10    further manipulation of polypeptide sequences or for the production of the encoded polypeptide in suitable host cells, either eukaryotic, such as yeast or plant cells, or prokaryotic, such as bacteria, for example *E. coli*. This can be achieved using recombinant DNA techniques and methodologies well known to those skilled in the art.

15                 Thus the present invention further provides recombinant nucleic acids comprising polynucleotide sequences which encode glycosyltransferases involved in the biosynthesis of lipopolysaccharides of bacteria of the genus *Helicobacter*, more particularly of lipopolysaccharides of the species *Helicobacter pylori* and various  
20    strains thereof. Most particularly, the invention provides recombinant nucleic acids comprising polynucleotides identical over their entire lengths to polynucleotides having sequences set out in Table 1, as well as fragments and variants of such sequences. Among fragments and variants, preferred are those coding for polypeptides retaining the biological function or activity of the reference  
25    polypeptides.

30                 The isolated polynucleotides and fragments thereof can also be used as DNA diagnostic probes specific to *H. pylori*, for diagnostic or similar purposes. They may be used, for example, to check whether or not the polynucleotides according to the present invention are transcribed in bacteria of an infected tissue. They may be also useful in diagnosis of the stage of infection and determining the specific pathogen involved.

The isolated polynucleotides of the present invention may further be used as hybridization probes for RNA, cDNA and genomic DNA, for example to isolate cDNA or genomic clones of other genes that have a high sequence similarity to the polynucleotides of the present invention. Such probes will comprise at least 5 15 bases, preferably at least 30 bases, but may have even more than 50 bases.

Preferred isolated or recombinant polypeptides of the present invention are those showing the activity of glycosyltransferases involved in biosynthesis of the bacterial lipopolysaccharides of bacteria of the genus *Helicobacter*, more particularly lipopolysaccharides of the species *Helicobacter pylori* and various strains thereof. Most particularly preferred are polypeptides coded by polynucleotides having sequences shown in Table 1 (SEQ ID NOs: 1, 3, 5, 7, 9, 11, 13 and 15), and also those which have at least 50% identity to polypeptides shown in Table 1 (SEQ ID NOs: 2, 4, 6, 8, 10, 12, 14 and 16), preferably at least 10 15 70% identity, more preferably at least 80% identity, most preferably at least 95% identity, polypeptides closely related thereto as well as fragments and variants thereof. Of the polypeptides having substantial identity to polypeptides of Table 1, preferred are those having the same biological function or activity as the polypeptides appearing in Table 1.

20 Polypeptides having amino acid sequences shown in Table 1 correspond to those coded by open reading frames HP0826 (SEQ ID NO: 2), HP0159 (SEQ ID NO: 4), HP0479 (SEQ ID NO: 6) and HP1191 (SEQ ID NO: 8) of the genomic DNA of *H. pylori* strain 26695, by open reading frames SS0826 (SEQ ID NO: 10), 25 SS0159 (SEQ ID NO: 12) and SS0479 (SEQ ID NO: 14) of the genomic DNA of *H. pylori* strain SS1, and by open reading frame PJ0479 of the genomic DNA of *H. pylori* strain PJ1. Among several others, these ORFs have been cloned and expressed in suitable host cells and their function has been determined *in vitro* using techniques well known to those skilled in the art and discussed more in 30 detail further in this disclosure.

Polypeptides of the present invention can be produced as discussed above in connection with recombinant nucleic acids of the present invention. They can be

recovered and purified from recombinant cell cultures by methods and techniques well known to those skilled in the art, including ammonium sulfate or ethanol precipitation, acid extraction, and various forms of chromatography, in particular ion exchange and high performance liquid chromatography (HPLC). Well known 5 techniques for refolding protein may be employed to regenerate active conformation when the polypeptide is denatured during isolation and/or purification.

The invention also relates to methods of screening compounds, to identify those 10 which enhance (agonists) or block (antagonists) the action of polynucleotides or polypeptides of the present invention. Of those, antagonists acting as bacteriostatic or bactericidal agents are of particular interest. Potential antagonists include small organic molecules, peptides, polypeptides and antibodies that bind to a polynucleotide or polypeptide of the present invention 15 and therefore inhibit its activity. Polynucleotides and polypeptides of the present invention may be used to assess the binding of small molecule substrates and ligands from various sources, including cells, cell-free preparations, chemical libraries, and natural product mixtures. The substrates and ligands may be natural substrates and ligands or may be structural or functional mimetics.

20 Polypeptides of the present invention are particularly useful for screening chemical compounds modulating the enzymatic activity of glycosyltransferases of *Helicobacter* origin involved in the biosynthesis of bacterial lipopolysaccharides, to identify those which enhance (agonists) or inhibit (antagonists or inhibitors) the 25 action of *Helicobacter* glycosyltransferases, in particular compounds that are bacteriostatic and/or bactericidal. The method of screening may involve high-throughput techniques and assays. In a typical assay, a synthetic reaction mix comprising a polypeptide of the present invention and a labelled substrate or ligand of such polypeptide is incubated in the absence and in the presence of a 30 candidate substance, a potential agonist or antagonist of the enzyme under study. This capability is reflected in decreased binding of the labeled ligand or in decreased production of a product from the labeled substrate. Detection of the rate or level of production of the product from the substrate may be enhanced by

using a suitable reporter system, such as a colorimetrically labelled substrate which is converted into a colorimetrically assayable product or a reporter gene responsive to changes in the enzymatic activity of the polypeptide.

- 5    The polypeptides of the present invention showing enzymatic activity of *Helicobacter* glycosyltransferases are also useful for the enzymatic synthesis of bacterial lipopolysaccharides and fragments thereof. When included in suitable reaction mixtures, these polypeptides catalyze the transfer of mono- or oligosaccharide residues to a suitable acceptor. In a preferred embodiment, the  
10    polypeptides of the present invention are used for the preparation of various mimics, analogues and derivatives of *Helicobacter* lipopolysaccharides.

In yet another aspect, the invention provides novel mutants of *Helicobacter* bacteria, in particular mutants of *H. pylori*, having mutated (deactivated) genes of  
15    glycosyltransferases involved in the biosynthesis of bacterial lipopolysaccharides, in particular of the core or O-chain regions of LPS. Structural analysis of LPS isolated from the mutants confirmed that O-chain synthesis has been affected by the mutations and revealed the exact structure of the truncated LPS molecules. The mutant strains were also shown to have a reduced capacity of gastric  
20    colonization.

The mutant bacteria expressing the truncated LPS and the LPS isolated from such mutants are useful as sources of antigens to be used in vaccination against *Helicobacter* bacteria, in particular against *H. pylori*. Such vaccines are normally  
25    prepared from dead bacterial cells, using methods well known to those skilled in the art, and usually contain various auxiliary components, such as an appropriate adjuvant and a delivery system. A delivery system aiming at mucosal delivery is preferred. Preferably but not essentially, the antigenic preparation is administered orally to the host, but parenteral administration is also possible. Live vaccines  
30    based on *H. pylori* mutants may also be prepared, but would normally require an appropriate vector for mucosal delivery. Vaccines of the present invention are useful in preventing and reducing the number of *H. pylori* infections and indirectly

in reducing the incidence of pathological conditions associated with such infections, in particular gastric cancer.

Chemically modified LPS isolated from mutants expressing the truncated LPS is  
5 a preferred antigen for use in vaccines according to the present invention. It is isolated from the bacteria and at least partially purified using techniques well known to those skilled in the art. Preparations of at least 70%, particularly 80%, more particularly 90%, most particularly 95% pure LPS are preferred. The purity of an LPS preparation is expressed as the weight percentage of the total  
10 Helicobacter antigens present in the preparation. The purified LPS can be used as antigen either directly or after being conjugated to a suitable carrier protein.

In the following, the invention will be described in still greater detail, by way of examples and with respect to the preferred embodiments.

15

#### **Identification and cloning of $\beta$ -1,4-galactosyltransferase**

A search of the *H. pylori* genomic database of translated proteins revealed three open reading frames (ORFs) (HP0826, HP0805 and HP0619) which exhibited  
20 limited homology with the *lex2B* gene from *Haemophilus influenzae* (39% identity) and the *lob1* gene from *Haemophilus somnus* (32% identity). While both the *lex2B* and *lob1* genes of *Haemophilus* have been shown to be involved in synthesis of the outer core region of the lipooligosaccharide (Jarosik *et al*, *Infect. Immun.* 62: 4861-4867 (1994); Inzana *et al*, *Infect. Immun.* 65: 4675-4681  
25 (1997)), to date no definitive function for either gene has been proposed. There is evidence that they are involved in addition of glucose (*lex2B*) and galactose (*lob1*) to the core heptose region. Both *lex2B* and *lob1* show significant homology to a larger group of LOS biosynthesis proteins which include the *H. influenzae* *lex1/lic2A* genes (Cope *et al*, *Mol. Microbiol.* 5: 1113-1124 (1994))  
30 and *lic2B* gene (High *et al*, *Mol. Microbiol.* 9: 1275 (1993)), *Neisseria lgtB* and *lgtE* genes (Wakarchuk *et al*, *J. Bio. Chem.* 271: 19166-19173 (1996)) and *lpsA* of *P. haemolytica* (Potter *et al*, *FEMS Microbiol. Lett.* 129: 75-81 (1995) which are all involved in outer core assembly. The *LgtB* and *LgtE* proteins of *N. meningitidis*

have been shown to be galactosyltransferases involved in the transfer of galactose in a  $\beta$ -1,4 linkage in the terminal lacto-N-neotetraose structure. LgtB is responsible for the addition of Gal to GlcNAc, an identical function to that described here for HP0826, while LgtE catalyses the addition of Gal to Glc 5 (Wakarchuk *et al, supra*). Clustal multiple sequence alignment of HP0826 amino acid (aa) sequence and lex2B, lob1 and lgtB aa sequences from this group of related LOS biosynthesis proteins did identify two regions of conservation spanning the regions in HP0826 from approx. aa90 to aa142 and aa189 to aa235 (see Fig 1). Limited homology was also observed with waaX from *E. coli* 10 (Heinrichs *et al, Mol. Microbiol.* 30: 221-232 (1998)), a putative core  $\beta$ -1,4-galactosyltransferase, only in the region spanning aa96-aa142 (data not shown). No significant homology was obtained with any putative glycosyltransferases involved in O-chain assembly from Gram-negative bacteria.

15 Synthetic oligonucleotide primers which contained BamHI restriction sites which flanked the 5' and 3' ends of HP0826, HP0619, and HP0805 respectively, were used in a PCR reactions containing chromosomal DNA of *H. pylori* 26695 or SS1 as a template. A single PCR product was obtained in each case and this was cloned into pUC19 to give plasmids pHp0826, pHp0805, and pHp0619. DNA sequencing was used to confirm the identity of the cloned PCR products from 20 26695 and SS1.

Three additional open reading frames of *H. pylori* genome, HP0159, HP1191 and HP0479, have been identified by BLAST analysis as potentially coding for LPS 25 glycosyltransferases. Of those, HP0159 displayed homology to the *rfaJ*, lipopolysaccharide 1,2-glucosyltransferase gene from a number of bacterial species, HP0479 and HP1191 displayed homology to *waaC* and *waaF* respectively, which are heptosyltransferase genes responsible for the addition of LD heptose to KDO in the core backbone.

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### **Functional analysis of lex2B homologues**

$\beta$ -1,4-galactosyltransferase activity has previously been detected in *H. pylori* (Chan *et al, Glycobiology* 5: 683-688 (1995)), but the gene(s) for this enzyme

have not been described. Enzyme activity was detected in extracts of *E. coli* pHP0826 but not from clones of HP0805 and HP0619 using the synthetic acceptor molecule FCHASE aminophenyl $\beta$ -GlcNAc and UDP-Gal as the donor. The lack of detectable activity in HP0805 and HP0619 clones could be a lack of 5 the appropriate donor/acceptor molecule for their respective enzymatic activities.  $\beta$ -1,4-galactosyltransferase activity was also present in parent *H. pylori* strains but not in the *H. pylori* HP0826 mutants. The assays were followed by TLC analysis of the reaction mixtures as previously described (Gilbert *et al*, *Eur. J. Biochem.* 249: 187-194 (1997)). A more sensitive capillary electrophoresis (CE) 10 analysis of the reaction mixtures clearly demonstrated a loss of galactosyltransferase activity in the mutants. The product of the enzymatic reaction had an identical CE mobility compared to a known FCHASE-aminophenyl- $\beta$ -N-acetyllactosamine standard, and was subjected to NMR analysis to determine the linkage. The  $^1\text{H}$  and  $^{13}\text{C}$  chemical shift data (Table 2) 15 and 1D NOE analysis were consistent with the linkage of the Gal being  $\beta$ -1,4 to the GlcNAc. The product was also sensitive to  $\beta$ -galactosidase.

**Table 2.** Linkage analysis of the product formed by HP0826 encoded protein.  
<sup>1</sup>H and <sup>13</sup>C chemical shifts of the glycoside of Gal- $\beta$ -1,4-GlcNAc- $\beta$ -FEX<sup>a</sup>

Residue	Position	H	C
<b>A, <math>\beta</math>-GlcNAc</b>	1	4.86	100.6
	2	3.91	55.8
	3	3.72	73.4
	4	3.72	79.0
	5	3.46	75.8
	6	3.74, 3.83	60.8
	NAc	1.91	22.9
<b>B, <math>\beta</math>-Gal</b>	1	4.46	103.8
	2	3.58	72.0
	3	3.68	73.4
	4	3.94	69.4
	5	3.73	76.3
	6	3.77, 3.77	62.0
FEXas		3.09	29.4
FEXms		2.80	36.9
FEXxs		3.57	37.6
FEXa1		6.92	118.2
FEXx1		7.28	124.4
FEXa2		7.17	132.5
EXm2		7.70	123.3
FEXx2		8.00	121.5
FEXa3		7.22	132.7
FEXa3'		7.13	131.1
FEXm3		6.82	121.5
FEXx3		6.91	104.3

5      <sup>a</sup> in ppm from the 600 MHz HSQC spectrum of the sample in D<sub>2</sub>O at 35°C. Chemical  
shifts are referenced to the methyl resonance of acetone set at 2.225 ppm for <sup>1</sup>H and  
31.07 ppm <sup>13</sup>C. The error is  $\pm$  0.03 ppm for <sup>1</sup>H and  $\pm$  0.3 for <sup>13</sup>C chemical shifts.  
The AMX spin system of CH<sub>2</sub>-CH<sub>2</sub>-S-CH<sub>2</sub> is at 3.09, 2.80, 3.57 ppm with J<sub>AM</sub>=6.4 Hz and  
with their respective <sup>13</sup>C signals at 29.4, 36.9 and 37.6 ppm. The aminophenyl A<sub>2</sub>X<sub>2</sub> spin  
10     system is at 6.92 and 7.28 ppm with J<sub>AX</sub>=8.7 Hz and their respective <sup>13</sup>C signals at 118.2  
and 124.4 ppm. The three AMX spin system for fluorescein carboxamido group with  
J<sub>AM</sub>=8-9 Hz and J<sub>MX</sub>= 1-2 Hz are at (7.17, 7.70, 8.00), (7.22, 6.82, 6.91) and (7.13, 6.82,  
6.91) ppm. Their respective <sup>13</sup>C signals are at (132.5, 123.3, 121.5), (132.7, 121.5,  
104.3) and (131.1, 121.5, 104.3) ppm.

**Functional analysis of rfaJ homologue (HP0159)**

Enzyme activity was detected in extracts of *E. coli* pHP0159 using the synthetic acceptor molecule FCHASE aminophenyl- $\alpha$ -maltose or FCHASE aminophenyl- $\alpha$ -glucose and UDP-Glc as the donor. Activity was also present in parent *H. pylori* strains but not in *H. pylori* HP0159 mutants. The assays were followed by TLC and CE analysis of the reaction mixtures as previously described (Gilbert *et al*, *Eur. J. Biochem.* 249: 187-194 (1997)). The more sensitive capillary electrophoresis (CE) analysis of the reaction mixtures demonstrated a loss of glucosyltransferase activity in the mutants. The product of the enzymatic reaction was subjected to NMR analysis to determine the linkage (Table 3). The  $^1\text{H}$  and  $^{13}\text{C}$  chemical shift data, and 1D NOE analysis were consistent with the linkage of Glc being  $\alpha$ -1,6 to the Glc.

**Table 3.** Linkage analysis of the product formed by HP0159 encoded protein.  
<sup>1</sup>H and <sup>13</sup>C chemical shifts of Glc- $\alpha$ -1,6-Glc- $\alpha$ -1,6-Glc- $\alpha$ -FEX<sup>a</sup>

Residue	Position	H	C
<b>A, <math>\alpha</math>-Glc-FEX</b>	1	5.35	98.3
	2	3.62	72.1
	3	3.80	74.1
	4	3.48	70.6
	5	3.72	72.1
	6	3.43, 3.69	66.5
<b>B, <math>\alpha</math>-Glc</b>	1	4.74	98.8
	2	3.47	72.2
	3	3.61	74.3
	4	3.48	70.6
	5	3.73	71.2
	6	3.59, 3.87	66.5
<b>C, <math>\alpha</math>-Glc (terminal)</b>	1	4.89	98.8
	2	3.52	72.5
	3	3.70	74.1
	4	3.41	70.5
	5	3.67	72.8
	6	3.74, 3.79	61.5
FEXas		3.02	29.3
FEXms		2.74	36.9
FEXxs		3.52	37.5
FEXa1		7.00	118.6
FEXx1		7.27	124.2
FEXa2		6.92	131.9
FEXm2		7.60	124.6
FEXx2		8.07	120.7
FEXa3		6.95	132.0
FEXa3'		6.92	131.9
FEXm3		6.69	119.6
FEXx3		6.79	104.1

5

<sup>a</sup> in ppm from the 600 MHz HSQC spectrum of the sample in D<sub>2</sub>O at 40°C. Chemical shifts are referenced to the methyl resonance of acetone set at 2.225 ppm for <sup>1</sup>H and 31.07 ppm for <sup>13</sup>C. The error is  $\pm$  0.03 ppm for <sup>1</sup>H and  $\pm$  0.3 for <sup>13</sup>C chemical shifts.

**Functional analysis of waaF homologue (HP1191)**

Complementation analysis was used to determine the function of the HP1191 from *Helicobacter pylori* strain 26695. The *H. pylori* HP1191 gene was amplified by PCR (see Table 6 for primer sequences used) and cloned into pUC19 to obtain pHp1191. WaaF mutant strain *S. typhimurium* 3789 was electroporated with this recombinant plasmid, and one of the resultant transformants selected for further study. SDS-PAGE was used to analyze LPS molecules produced by the relevant *S. typhimurium* strains. The LPS of the wild type strain formed the ladder like pattern indicative of the presence of the O antigen repeat unit whereas the LPS of the *S. typhimurium* waaF mutant appeared as a single fast migrating band. The migration pattern of this mutant was not affected by the presence of the plasmid vector. However, when the *H. pylori* gene HP1191 was present *in trans* in strain 3789, this *S. typhimurium* mutant synthesized an LPS which migrated in a pattern identical to that obtained with the LPS of the wild type strain. This confirmed the activity of HP1191 to be involved in catalyzing the addition of a second heptose molecule onto the heptose linked directly to KDO in the core.

**Construction of *H. pylori* mutants carrying a disrupted HP0826 gene**

In order to determine the role of the HP0826 ORF in LPS biosynthesis, *H. pylori* mutants carrying a disrupted HP0826 gene were constructed by allelic exchange. Briefly, the HP0826 ORF cloned in pUC19 was disrupted by using reverse primers 5'TACAGATCGCTTCATTGAGTTCT3' and 5'CCAAGAGTTAGGCTATATCCGCTT3' in a PCR reaction and ligating a kanamycin resistance cassette (or Km<sup>r</sup>) to the gel purified product to make plasmid pHp0826::kan. *H. pylori* strains 26695, NCTC11637, O:3 and Sydney strain (SS1) were transformed with plasmid pHp0826::kan DNA following the procedure of Haas *et al*, *Mol. Microbiol.* 8:753-760 (1993). This construct contains 515bp of homologous DNA upstream of the mutation and 464bp downstream of the mutation. Following transformation, cells were plated on blood agar containing kanamycin (20 µg/ml). Km<sup>r</sup> colonies were isolated and passaged on the same medium. Individual colonies were selected and screened for the presence of a double cross over mutation in the chromosome of the kan mutant.

To assess the insertion site of the disrupted gene PCR analysis was used, with chromosomal DNA from parent and mutant *H. pylori* strains as templates and the primer pair 5'ACACTGGCATCATAAAAT3' and 5'CCATGCGAAGTTATGAGCT3' which are internal in the structural gene. This analysis demonstrated conclusively that the Km<sup>r</sup> cassette was inserted into the chromosomal copy of HP0826. The primer pair amplified the expected 212bp fragment in the parent strain, but resulted in a 1.6kb fragment consistent with insertion of the 1.4kb Km<sup>r</sup> cassette. Plasmid vector sequences were not detected by Southern blotting and a single 1.7kb Hind III fragment corresponding to insertion of the kan cassette in the HP0826 ORF was present in chromosomal DNA's of 26695::0826kan mutant and SS1::0826kan mutant but not in parental DNA when probed with the kan cassette. These data confirm that the insertion mutant was the result of a double cross-over event. Four kanamycin resistant transformants were independently tested to verify that gene disruption and gene replacement had occurred. All four mutants grew normally *in vitro* (as assessed by OD vs viable numbers) and produced a truncated LPS as assessed by electrophoretic mobility on SDS-PAGE gels. The overall protein composition of the total membrane fraction was unchanged in the knockout mutants of SS1 and 26695 as assessed by SDS-PAGE and Coomassie blue staining. The contribution of polar effects to the phenotype of the HP0826 mutant is highly unlikely as a transcriptional terminator lies immediately downstream of the HP0826 ORF, the transcriptional organization switches strands and the downstream annotated ORF HP0827 is unrelated to LPS biosynthesis.

The construction of *H. pylori* mutants carrying disrupted HP0159 and HP0479 genes was carried out in essentially the same manner as above.

#### **Genomic Organization and Allelic Variation of SS1**

To ascertain if the structural organization found in 26695 and J99 is conserved within the SS1 genome, PCR amplification and sequencing of the HP0826 homologue and flanking sequence was obtained from SS1. As with 26695 and J99, the upstream and downstream ORFs are conserved although variation in the intervening sequence was evident. Allelic variation of SS1 HP0826 resulted

in 31 base pair differences between SS1 and 26695 and 46 base pair differences between SS1 and J99. These differences in DNA sequence results in a total of 9 amino acid changes in the SS1 protein when compared with 26695 and J99 amino acid sequences. In both comparisons the variations were located 5 predominately at the N and C terminal region of the protein.

#### **SDS-PAGE analysis of *H. pylori* HP0826 mutants**

To characterize the effect of the HP0826 mutation on LPS structure in *H. pylori*, proteinase K digested whole cell lysates from both parent and mutant cells grown 10 in broth were analyzed by SDS-PAGE. Silver staining revealed significant differences in the electrophoretic mobility of LPS isolated from parent and mutant cells of each strain examined. LPS from strains 26695, SS1, O:3 and NCTC11637 appeared to have typical high molecular weight, smooth form LPS (S-LPS), while the HP0826 mutant of each strain no longer produced the S-LPS, 15 but appeared to produce a semi-rough type LPS. Immunoblotting with monoclonal antibodies to Lewis X ( $Le^X$ ) and Lewis Y ( $Le^Y$ ) antigens confirmed that the LPS from all mutants no longer displayed immunoreactive material of high molecular weight typical of the corresponding parental O-chain which displays Lewis antigens.

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#### **SDS-PAGE analysis of *H. pylori* HP0159, 0479 and 1191 mutants**

To characterize the effect of the HP0159, 0479 and 1191 mutations on LPS structure in *H. pylori*, proteinase K digested whole cell lysates from both parent and mutant cells grown in broth were analyzed by SDS-PAGE. Silver staining 25 revealed significant differences in the electrophoretic mobility of LPS isolated from parent and mutant cells of each strain examined. In all cases, LPS from mutant cells no longer produced S-type LPS but instead only a fast migrating rough type LPS was observed.

#### **30 Structural investigations of *H. pylori* HP0826 LPS mutants of strains 26695, SS1, and NCTC 11637**

The LPS molecules of *H. pylori* strains 26695, SS1 ( M. A. Monteiro *et al*, *Eur. J. Biochem.* 267: 305-320 (2000) and type strain NCTC 11637 (Aspinall *et al*,

supra) have been determined to carry O-chain regions that express Le<sup>x</sup> and Le<sup>y</sup> blood-group determinants. These Lewis-mimicking O chains were shown to be covalently connected to a core oligosaccharide. Sugar composition analysis (Table 4) of the intact LPSs of *H. pylori* 26695::HP0826kan, SS1::HP0826kan and NCTC 11637::HP0826kan demonstrated a clear reduction in levels of those sugars known to form the O chain components, namely L-Fuc, D-Gal and D-GlcNAc, when compared to parent LPSs.

10 Table 4. Approximate molar ratios of the alditol acetate derivatives of 26695, SS1 and NCTC 11637 HP0826 isogenic mutants intact LPSs. Numbers in parentheses indicate ratios obtained for respective parent strains. Analyses performed on LPS prepared from broth grown cells.

Strain	L-Fuc	D-Glc	D-Gal	GlcNAc	DD-Hep	LD-Hep
15 26695::Hp0826kan	0.8 (6)	6 (7)	1 (10)	1 (8)	2 (2)	1.8 (1.6)
SS1::Hp0826kan	0.8 (6)	2 (2)	1 (10)	1 (8)	2 (2)	1.8 (1.6)
NCTC11637::Hp0826kan	0.8 (6)	6 (7)	1 (10)	1 (8)	2 (2)	1.8 (1.6)

20 Methylation linkage analysis performed on the intact *H. pylori* mutant LPSs from each strain showed the presence of terminal and 3-substituted Fuc, terminal, 3-, and 6-(except in SS1 strain) substituted Glc, terminal, 3- and 4-substituted Gal, 2-(only in 26695), 3-(only in 26695), 6-(only in 26695), 7- and 2,7-substituted DD-Hep, 2- and 3,7-substituted LD-Hep, and terminal and 3-substituted GlcNAc units. All sugars were present in the pyranose conformation. In order to obtain sugar sequence information of the outer-extremities of the LPS molecule (O-chain perimeter), a fast atom bombardment-mass spectrometry (FAB-MS) experiment in the positive ion mode was carried out on the methylated intact mutant LPSs from each strain. The FAB-MS spectra showed several A-type primary glycosyl oxonium ions of defined composition. The trace amounts of terminal GlcNAc that were observed in the linkage analyses were also detected in each of the three mutant LPS FAB-MS spectra at m/z 260 [GlcNAc]<sup>+</sup> (Fig. 2). A-type primary glycosyl oxonium ions containing Lewis blood-group related Fuc

and GlcNAc residues were observed at m/z 434 [Fuc, GlcNAc]<sup>+</sup>, 508 [GlcNAc, Hep]<sup>+</sup>, and 682 [Fuc, GlcNAc, Hep]<sup>+</sup>. The ion m/z 434 stood for a disaccharide composed of Fuc and GlcNAc and ion m/z 508 characterized a possible connection between the O-chain related GlcNAc and a heptose from the core region. The ion m/z 682 [Fuc, GlcNAc, Hep]<sup>+</sup> represented a moiety containing GlcNAc and Fuc residues of the O-chain region and a single heptose unit from the core region which bridges the O-chain and the core oligosaccharide. Since no terminal Hep unit was detected, these m/z 508 and 682 ions must originate from cleavage at the heptose glycosidic bond and represent a partial O-chain repeating unit [Fuc, GlcNAc, Hep]<sup>+</sup>. No 3,4-substituted GlcNAc, 2-substituted Gal and no m/z 638 (characteristic of Le<sup>x</sup>) and 812 (characteristic of Le<sup>y</sup>) glycosyl oxonium ions were detected, and therefore no evidence of an O-chain containing Le<sup>x</sup> or Le<sup>y</sup> determinants was found in these analyses of 26695::HP0826kan, SS1::HP0826kan and NCTC 11637::HP0826kan LPSs. In addition, higher mass ions in the FAB-MS spectrum of NCTC11637::HP0826kan at m/z 886 [Fuc, GlcNAc, Hep, Glc]<sup>+</sup>, 1090 [Fuc, GlcNAc, Hep, Glc<sub>2</sub>]<sup>+</sup>, and 1294 [Fuc, GlcNAc, Hep, Glc<sub>3</sub>]<sup>+</sup> (Fig. 2) represented the characteristic glucosylated by a [(1-6)- $\alpha$ -glucan] heptose unit (Aspinall *et al*, *supra*) in strain NCTC 11637 and 26695 (Fig. 2). The same primary ions were also observed in the FAB-MS spectrum of the methylated LPS of 26695::HP0826kan, but not of SS1::HP0826kan, in line with the structural findings in the parent strains (M. A. Monteiro, unpublished). In the three FAB-MS spectra, the primary ion m/z 668 and its corresponding secondary ion m/z 228 (Fig. 2) pointed to the presence of the type 1 linear B blood-group [Gal-(1-3)-Gal-(1-3)-GlcNAc] antigen, a blood-group determinant found in the LPSs of 26695, SS1 (M. A. Monteiro, unpublished), and in NCTC 11637 (Monteiro *et al*, *J. Biol. Chem.* 273: 11533-11543 (1998)). The glycose units emanating from the core oligosaccharide regions were of the same type as those found in the respective parent LPSs. The GlcNAc and Fuc units observed were remnants of an incomplete O chain. A comparison of the structures identified in parent and mutant LPS from 26695 and SS1 and the respective HP0826,0159 and 0479 isogenic mutants is presented in Fig 3.

**Structural characterization of *H. pylori* LPS mutants 26695::HP0159kan and SS1:: HP0159kan**

Growth of bacterial strains and isolation of LPS by hot aqueous phenol method  
5 were carried out as described previously (Logan et al, *Mol. Microbiol.* 35: 1156-1167 (2000)). Sugar analysis of the intact LPS of *H. pylori* 26695:: HP0159kan, SS1:: HP0159kan, O:3:: HP0159kan showed significant reduction in L-Fuc, D-Gal, and DD-Hep (for serotype O:3 mutant) when compared with the parent LPS indicating the presence of the structure devoid of O-chain and DD-heptan.  
10 Methylation analysis of the intact LPS from each strain showed the presence of terminal and 3-substituted L-Fuc, terminal and 4-substituted D-Glc, terminal, 3- and 4-substituted D-Gal, terminal, 2-, 6-, 7- and 2,7-substituted DD-Hep, terminal, 2- and 3-substituted LD-Hep and terminal, 3-substituted and 4-substituted D-GlcNAc. All sugars were present in the pyranose form. In addition, methylation  
15 analysis of LPS from 26695::HP0159kan and O:3::HP0159kan revealed the presence of 4-substituted D-Glc, no 6-substituted D-Glc was observed. NMR analysis of a high molecular mass fraction, isolated by gel filtration chromatography from a partially delipidated LPS (1.5% acetic acid, 1h, 100°C) from 26695:: HP0159kan by gel filtration chromatography, indicated it to contain  
20 β-1,4-linked glucan, a contaminant produced by some strains of *H. pylori* (Knirel et al, *Eur. J. Biochem.* 266: 123-131 (2000)). In order to deduce the sequence information on the outer extremities of the LPS molecule, permethylated intact LPS from each strain was subjected to the fast-atom-bombardment mass spectrometric analysis in the positive mode. A-type primary glycosyl oxonium  
25 ions containing Lewis blood group related Fuc and GlcNAc residues were observed at m/z 260 [GlcNAc]<sup>+</sup> and m/z 682 [Fuc,GlcNAc, Hep]<sup>+</sup>. No higher mass ions representing a glucosylated DD-heptose unit were detected. This evidence together with the absence of 6-substituted glucose in methylation analysis indicated this LPS mutant to be deficient in the biosynthesis of α(1-6)-glucan  
30 present in both 26695 and O:3 parent strains. Absence of the 3-substituted glucose in methylation analysis of LPS from 26695::HP0159kan, SS::HP0159kan, suggested that addition of a 1,3-linked glucopyranosyl residue was also impaired by this mutation. In the three FAB-MS spectra, the primary ion m/z 668 and its corresponding secondary ion m/z 228 pointed to the presence of

the type 1 linear B blood group [Gal(1-3)Gal(1-3)GlcNAc] antigen, a blood group antigen found in the LPS of 26695 and SS1 (Monteiro *et al*, *Eur. J. Biochem.* 267:305-320 (2000)). Other Lewis blood group-related secondary ions were observed at m/z 228 (260-32) [GlcNAc]<sup>+</sup>, 402 (434-32) [Fuc,GlcNAc]<sup>+</sup>, 576 (608-32) [Fuc (1-3)Fuc (1-4)GlcNAc]<sup>+</sup> as previously described (Monteiro *et al*, *J. Biol. Chem.* 273: 11533-11543 (1998), Logan *et al*, *Mol. Microbiol.* 35: 1156-1167 (2000)).

LPS from 26695::HP0159kan was treated with 0.1 M sodium acetate buffer, pH 10 4.2 (2 h, 100°C) and following the removal of lipid A by low speed centrifugation, subjected to the gel filtration chromatography on a Bio-Gel P-2 column, followed by capillary electrophoresis-electrospray mass spectrometry (CE-ESMS) as described previously (Thibault and Richards, *Meth. Mol. Biol.* 145: 327-343 (2000)). The CE-ESMS spectrum of the delipidated LPS confirmed the presence 15 of a major glycoform produced by the 26695::HP0159 mutant LPS, corresponding to FucGlcNAcHex<sub>2</sub>Hep<sub>4</sub>(PE)KDO (m/z 902, doubly protonated ion). MS-MS of the doubly charged ion (m/z 902) (Fig. 4) afforded a singly charged fragment at m/z 1601 consistent with the loss of GlcNAc (and its anhydro form at m/z 1583) which subsequently lost Fuc and Hep residues to 20 afford a fragment ion at m/z 1262. A comparison of the structures identified in parent and HP0159 mutant LPS is presented in Fig. 3.

**Structural characterization of *H. pylori* LPS mutants 26695::HP0479kan and SS1::HP0479kan.**

Sugar analysis of the HP0479 LPS mutants indicated reduction in the amount of 25 L-Fuc, D-Gal and DD-Hep and methylation analysis confirmed this. Methylation analysis of the intact LPS from each strain indicated absence of 3-substituted and 6-substituted D-Glc, 3-substituted DD-Hep and 6-substituted DD-Hep (for 30 O:3::HP0479 and 26695::HP0479 LPS) and a significant decrease in 2-substituted DD-Hep, suggesting deficiencies in the core biosynthesis.

FAB-MS analysis in the positive mode of the permethylated LPS from each strain indicated the presence of primary glycosyl oxonium ions at m/z 260 [GlcNAc]<sup>+</sup>

and m/z 434 [Fuc,GlcNAc]<sup>+</sup> and secondary glycosyl oxonium ions at m/z 228 (260-32) [GlcNAc]<sup>+</sup> and m/z 402 (434-32) [Fuc,GlcNAc]<sup>+</sup>. This evidence together with the absence of the primary glycosyl oxonium ion at m/z 682 [Fuc, GlcNAc, Hep]<sup>+</sup> suggested that the mutant LPS structure was lacking DD-Hep residue  
5 which bridges O-chain and the core oligosaccharide in the respective parent LPS (Monteiro *et al*, *Eur. J. Biochem.* 267: 305-320 (2000), Logan *et al*, *Mol. Microbiol.* 35: 1168-1179 (2000)). LPS from SS1:: HP0479 and 26695 was delipidated and desaltsed following gel filtration chromatography on a Bio-Gel P-2 column. Fractions containing core oligosaccharide components were subjected to  
10 the mass spectrometric analysis using combined capillary zone electrophoresis-electrospray-mass spectrometry (CZE-ESMS) in the positive mode, followed by MS/MS analysis of the most abundant oligosaccharide fragments. The product ion spectrum showed two major singly charged fragment ions at m/z 1612 and m/z 1246, containing an anhydro-KDO. The fragment ion at m/z 1612 could be  
15 assigned to the glycoform FucGlcNAcHex<sub>2</sub>Hep<sub>3</sub>(PE)KDO (Fig. 5), based on the linkage and FAB-MS analyses data and recent structural studies (Monteiro *et al*, *Eur. J. Biochem.* 267: 305-320 (2000)). The MS/MS spectrum of m/z 1246 was consistent with the core fragment Hex<sub>2</sub>Hep<sub>3</sub>(PE)KDO as confirmed by a consecutive cleavage of glycosidic bonds yielding a direct sequence assignment.  
20 These structural assignments are consistent with the presence of 2,7-substituted DD-Hep, 7-substituted DD-Hep and 2-substituted DD-Hep in the methylation analysis of LPS mutants 26695::HP0479kan, SS1::HP0479kan, O:3::HP0479kan. Absence of the first DD-Hep which serves as a link between the O-chain and the  
25 core oligosaccharide and is glycosylated by 1,6-glucan, resulted in the loss of O-chain and DD-heptan (for serotype O:3). A comparison of the structures identified in parent and HP0479 mutant LPS is presented in Fig. 3.

### Mouse Colonization Studies

The role of S-type LPS in gastric colonization was investigated using the SS1 strain of *H. pylori* which others (Lee *et al*, *Gastroenterology* 112: 1386-1397 (1997); Ferrero *et al*, *Infect. Immun.* 66: 1349-1355 (1998); Conlan *et al*, *Can. J. Microbiol.* 45:975-980 (1999)) have shown to be capable of colonising the stomachs of mice, including the CD1 strain used in the present study. Both  
30

parental SS1 and SS1 HP0826 mutant which was obtained by natural transformation were used to orogastrically inoculate mice. The parent SS1 cells produce considerable amounts of S type LPS displaying Lewis Y epitopes while cells in which HP0826 has been inactivated produce a faster migrating, rough 5 type LPS molecule no longer displaying Lewis epitopes. To minimise the likelihood that any observed differences in *in vivo* behaviour arose as a result of exogenous influences, care was taken to ensure that the mutant and parental strains underwent equivalent *in vitro* manipulations before being gavaged into mice. In an initial experiment, groups of mice were gavaged with either wild-type 10 or mutated *H. pylori* SS1. Representative mice from each group were killed 6 or 12 weeks later and the stomach burdens of *H. pylori*, and level of *Helicobacter*-specific circulating immunoglobulin G determined. By 6 weeks of infection,  $5.65 \pm 0.26 \log_{10}$ CFU (colony-forming units) of wild-type bacteria were recovered from the stomachs of mice ( $n=4$ ) challenged with this organism, whereas only 15  $4.27 \pm 0.26 \log_{10}$ CFU of the mutant bacteria were recovered from the stomachs of mice gavaged with it. This 24-fold decreased recovery of mutant *versus* wild-type *H. pylori* SS1 was statistically significant according to the Mann-Whitney Rank Sum Test ( $p<0.05$ ). Similarly, by 12 weeks there was a 10-fold difference 20 in numbers of wild-type ( $5.81 \pm 0.51 \log_{10}$ CFU,  $n=5$ ) and mutant ( $4.79 \pm 0.43 \log_{10}$ CFU,  $n=5$ ) bacteria recovered, and this too was statistically significant ( $p<0.05$ ). PCR performed on digested stomach tissue confirmed the above findings, indicating that the decreased recovery was not due to any innate unculturability of the mutant bacteria. Likewise, by 12 weeks of infection sera 25 from mice infected with wild-type SS1 all reacted by ELISA against a sonicate of *H. pylori* as coating antigen (average IgG titre =  $1270 \pm 2166$ ) whereas only 3/5 mice infected with mutant SS1 had seroconverted (mean IgG titre of seropositives =  $123 \pm 94$ ). Additionally, when either parental or mutant LPS was used as the coating antigen in ELISA, only mice infected with the parental strain of *H. pylori* showed evidence of seroconversion.

30

To determine whether the colonisation differences observed in the aforementioned experiment were due to an initial inability of the mutant strain to colonise or due to its subsequent elimination, a complementary experiment

examined gastric colonization levels of parental and mutated *H. pylori* SS1 at 1 and 3 weeks post-challenge. By one-week post-challenge,  $5.81+/-0.29 \log_{10}\text{CFU}$  ( $n=5$ ) of wild-type bacteria, but only  $3.94+/-0.33 \log_{10}\text{CFU}$  ( $n=5$ ) of the mutant bacteria were recovered from the stomachs of the respectively infected mice.

5 This 74-fold difference was statistically significant ( $P< 0.05$ ) and convincingly shows that *H. pylori* SS1 bacteria unable to produce S-type LPS are significantly impaired in their ability to initially colonise the murine stomach. In this experiment, approximately 17-fold more wild-type than mutant *H. pylori* ( $5.4+/-0.34 \log_{10} \text{CFU}$ ,  $n=5$  versus  $4.18+/-0.14 \log_{10}\text{CFU}$ ,  $n=5$ ) were recovered from the stomachs of

10 relevant mice at three weeks of infection.

Results of mouse colonization experiments for the parent (SS1) strain of *H. pylori* and their mutant strains SS0826, SS0159 and SS0479 are summarized in Table 5.

15

**Table 5.** Mouse colonization data. Numbers in the table show levels of colonization of mice stomachs (as  $\log_{10}\text{CFU}/\text{stomach} +/- \text{ standard deviation}$ ) after the indicated number of weeks (WK) of infection. ND: not determined BDL: less than 500 bacteria

20

	STRAIN	WK 1	WK 3	WK 6	WK 12	WK 20
<b>EXP 1</b>	<b>SS1</b>	$5.81 +/- 0.29$ (n = 5)	$5.40 +/- 0.34$ (n = 5)	$5.65 +/- 0.26$ (n = 4)	$5.81 +/- 0.51$ (n = 5)	ND
	<b>SS0826</b>	$3.94 +/- 0.33$ (n = 5)	$4.18 +/- 0.17$ (n = 5)	$4.27 +/- 0.26$ (n = 4)	$4.79 +/- 0.43$ (n = 5)	ND
<b>EXP 2</b>	<b>SS1</b>	$5.43 +/- 0.03$ (n = 4)	ND	ND	$5.94 +/- 0.33$ (n = 5)	$5.84 +/- 1.10$ (n = 5)
	<b>SS0159</b>	$3.37 +/- 0.20$ (n = 4)	ND	ND	$3.09 +/- 0.42$ (n = 5)	< 3.76 (n = 5)
<b>EXP 3</b>	<b>SS1</b>	$4.76 +/- 0.93$ (n = 5)	ND	ND	$5.02 +/- 1.06$ (n = 5)	ND
	<b>SS0479</b>	BDL (n = 5)	ND	ND	BDL (n = 5)	ND

- Exp 1: Individual mice inoculated by gavage on D1, D3, D6 with 0.2ml of broth grown cells suspended in PBS at cell concentration of ~1 x 10<sup>10</sup>/ml.
- 5    Exp 2: Individual mice inoculated by gavage on D1 + D3 with 0.2ml of broth grown cells suspended in PBS at cell concentration of ~2 x 10<sup>10</sup>/ml.
- 10    Exp 3: Individual mice inoculated by gavage on D1 and D3 with 0.2ml of broth grown cells suspended in PBS at cell concentration of 4.7x10<sup>10</sup>/ml (D1) and 1x10<sup>7</sup>/ml (D3)

15    The above data show that all the mutants with disrupted genes have a reduced ability to colonize the murine stomach, as compared with the parent strain. SS0479 strain (*H. pylori* strain SS1 having disrupted gene HP0479) is the least capable of colonization.

20    **EXPERIMENTAL**

**Bacterial strains and culture conditions**

*Helicobacter pylori* strain 26695 (Tomb *et al, supra*) used for the initial cloning was obtained from R. A. Alm, Astra, Boston. *H. pylori* strain SS1 was obtained from A. Lee. *H. pylori* reference strain ATCC43504 and *H. pylori* serogroup O:3 isolate were from J. Penner. PJ1 was a fresh clinical isolate of *H. pylori*. Helicobacter strains were grown on at 37°C on antibiotic supplemented (Lee *et al, supra*) trypticase soy agar plates containing 7% horse blood (GSS agar) in a microaerophilic environment for 48h (Kan 20 µg/ml). For growth in liquid culture, antibiotic supplemented Brucella broth containing 5% fetal bovine serum, was inoculated with *H. pylori* cells harvested from 48h trypticase soy agar/horse blood plates and incubated for 36h in a Trigas (Nuair, Plymouth, MN) incubator (85% N<sub>2</sub>, 10%CO<sub>2</sub>, 5%O<sub>2</sub>) on a shaking platform. *Escherichia coli* strain DH5 $\alpha$  was used as host for plasmid cloning experiments and was grown on L-agar plates at 37°C supplemented with ampicillin (50µgml<sup>-1</sup>) and/or kanamycin (20µgml<sup>-1</sup>)

**β-1,4-galactosyltransferase activity**

Glycosyltransferase assays were performed essentially as described previously (Gilbert *et al.*, *supra*). Cells were scraped from a 3 day old plate culture of *H. pylori*, the cells were stored frozen at -20°C. Cell extracts were made by mixing 5 the cell pellet with 2 volumes of glass beads, and grinding with a ground glass pestle in the microcentrifuge tube. The paste was extracted twice with 50 µl of 50 mM MOPS-NaOH buffer pH 7.0. Reactions contained 0.5 mM FCHASE-aminophenyl-β-GlcNAc, 10 mM MnCl<sub>2</sub>, 0.5 mM UDP-Gal, 50 mM MOPS-NaOH pH 7.0, and 10 µl of cell extract in a final volume of 20 µl. For reactions with the 10 cell extracts of *H. pylori* the reactions were incubated 3-5 h at 37°C, whereas with the extracts containing the recombinant enzyme the reactions times were 30 – 60 min at 37°C. The TLC and CE analysis was performed as previously described (Gilbert *et al.*, *supra*). For TLC analysis 0.5 µl of the reaction mixture were spotted and developed and for CE analysis samples were diluted to an 15 FCHASE-aminophenyl-β-GlcNAc concentration of 10 µM prior to analysis.

**Recombinant DNA techniques and nucleotide sequence analysis**

DNA sequencing of PCR products was performed using an Applied Biosystems (model 370A) automated DNA sequencer using the manufacturers cycle 20 sequencing kit. All standard methods of DNA manipulation were performed according to the protocols of Sambrook *et al*, *Molecular Cloning: A Laboratory Manual*, 2<sup>nd</sup> edn. Cold Spring Harbor, New York: Cold Spring Harbor Laboratory Press (1989). DNA probes for Southern blotting were labelled with DIG-11-dUTP using DIG-High Prime (Boehringer Mannheim, Montreal, Canada) and detection 25 of hybridized probe with DIG Luminescent Detection Kit (Boehringer Mannheim Montreal, Canada). Primers used for the PCR gene amplification and for mutant constructs are shown in Table 6.

Table 6. Primer sequences for PCR amplification of HP0826, HP0159, HP0479 and HP1191 genes and for construction of respective mutant strains .

	<u>Primer</u>	<u>Primer (5'-3') sequence</u>
5	HP0826-F1	cggatccGGTTTTATAGCCATGATGC
	HP0826-R1	cggatccAAGGCGGTTAACGTTTGTTC
	HP0826-mut1	TACAGATCGCTTCATTGAGTTCT
	HP0826-mut2	CCAAGAGTTAGGCTATATCCGCTT
10	HP0159-F1	cggatccTGTCAAATTGCCCTATAGCGT
	HP0159-R1	cggatccACCTATTTAGGGAAACCGCT
	HP0159-mut1	GCCGGGTTTTAGTCGTGAAT
	HP0159-mut2	AGGGAAAAGGCTTGACGAGG
15	HP0479-F1	GCCTTTATCAAGCTAGAG
	HP0479-R1	CATAAATGTCCTAACAAAGC
	HP0479-mutF1	CAAACCCGCCAGGAGTTG
	HP0479-mutR1	GGTTATGGGAATGAATTGG
20	HP1191-F1	cggatccCGGTCTTAAACCCGCTCAACA
	HP1191-R1	cggatccCCGCTCTCACGCCTTAA

#### **Site specific mutagenesis of HP0826**

25 HP0826 clone of *Helicobacter pylori* strain 26695 was mutagenized in *E. coli* by ligation of the Km<sup>r</sup> cassette described by Labigne et al (*J. Bacteriol.* 170: 1704-1708 (1988)) to pUC19 containing the HP0826 gene. Deletion of a central 66bp region of the gene was achieved by reverse PCR (Pwo polymerase, Boehringer Mannheim) using the outward primers 5'TACAGATCGCTTCATTGAGTTCT3' and 30 5'CCAAGAGTTAGGCTATATCCGCTT3' followed by blunt end ligation with the Km<sup>r</sup> cassette. The mutated allele was returned to Helicobacter by natural transformation according to the method of Haas et al (*supra*).

#### **Electrophoresis and Western blotting**

35 SDS-PAGE was performed with a mini-slab gel apparatus (Biorad) by the method of Laemmli (*Nature* 227: 680-685 (1970)). LPS samples were prepared from whole cells according to a previously described method (Logan et al, *Infect. Immun.* 45: 210-216 (1984)), equivalent amounts loaded in each lane and stained according to Tsai et al (*Anal. Biochem.* 119: 115-119 (1982)) or 40 transferred to nitrocellulose for immunological detection as previously described

(Logan *et al, supra*). Anti Lewis monoclonal antibodies (Signet Laboratories Inc, Dedham, MA) were used at 1:500 dilution.

#### **Isolation of membrane fraction**

5 Broth grown cells (18h) were harvested and resuspended in 20mM Tris (pH 7.4). Following sonication (3x60sec) intact cells were removed by centrifugation at 4000xg, and membranes sedimented by centrifugation at 40,000xg, washed in 20mM Tris (pH7.4) re-centrifuged, and resuspended in 0.5ml 20mM Tris (pH7.4). Equivalent amounts of SS1, 26695 parent and mutant strains were analyzed by  
10 SDS-PAGE and stained by Coomassie Blue.

#### **Isolation of Lipopolysaccharides**

The LPSs were isolated by the hot phenol-water extraction procedure (Westphal *et al, Meth. Carbohydr. Chem.* 5: 83-91 (1965)). The LPSs were purified by gel-  
15 permeation-chromatography on a column of Bio-Gel P-2 (1m x 1cm) with water as eluent. In all cases, only one carbohydrate positive fraction was obtained which eluted in the high  $M_r$  range (Dubois *et al, Anal. Chem.* 28: 350-356 (1956)). These intact *H. pylori* LPSs then were used for chemical analyses.

20 **Sugar Composition and Methylation Linkage Analyses**

Sugar composition analysis was performed by the alditol acetate method (Sawardeker *et al, Anal. Chem.* 39:1602-1604 (1967)). The hydrolysis was done in 4M trifluoroacetic acid at 100°C for 4h or 2M trifluoroacetic acid at 100°C for 16h followed by reduction in H<sub>2</sub>O with NaBD<sub>4</sub>, and subsequent acetylation with acetic anhydride and with residual sodium acetate as the catalyst. Alditol acetate derivatives were analyzed by gas-liquid-chromatography mass-spectrometry (GLC-MS) using a Hewlett-Packard chromatograph equipped with a 30 m DB-17 capillary column [210°C (30 min) to 240°C at 2°C/min] and MS in the electron impact (EI) mode was recorded using a Varian Saturn II mass spectrometer.  
25 Methylation linkage analysis was carried out by the NaOH/DMSO/CH<sub>3</sub>I procedure (Ciucanu *et al, Carbohydr. Res.* 131: 209-217 (1984)) and with characterization of permethylated alditol acetate derivatives by GLC-MS in the EI mode (DB-17 column, isothermally at 190°C for 60 min).

**Fast Atom Bombardment-Mass Spectrometry (FAB-MS)**

A fraction of the methylated sample was used for positive ion fast atom bombardment-mass spectrometry (FAB-MS) which was performed on a Jeol 5 JMS-AX505H mass spectrometer with glycerol(1) : thioglycerol(3) as the matrix. A 6 kV Xenon beam was used to produce pseudo molecular ions which were then accelerated to 3kV and their mass analyzed. Product ion scan (B/E) and precursor ion scan ( $B^2/E$ ) were preformed on metastable ions created in the first free field with a source pressure of  $5 \times 10^{-5}$  torr. The interpretations of positive ion 10 mass spectra of the permethylated LPS derivatives were as previously described by Dell *et al* (Carbohydr. Res. 200: 59-67 (1990)).

**Electrospray mass spectrometry**

Samples were analyzed on a crystal Model 310 CE instrument (ATI Unicam, 15 Boston, MA, USA) coupled to an API 3000 mass spectrometer (Perkin-Elmer/Sciex, Concord, Canada) via a microlonspray interface. A sheath solution (isopropanol-methanol, 2:1) was delivered at a flow rate of 1  $\mu\text{L}/\text{min}$  to a low dead volume tee (250  $\mu\text{m}$  i.d., Chromatographic Specialties, Brockville, Canada). All aqueous solutions were filtered through a 0.45- $\mu\text{m}$  filter (Millipore, Bedford, MA, 20 USA) before use. An electrospray stainless steel needle (27 gauge) was butted against the low dead volume tee and enabled the delivery of the sheath solution to the end of the capillary column. The separation were obtained on about 90 cm length bare fused-silica capillary using 10 mM ammonium acetate/ammonium hydroxide in deionized waster, pH 9.0, containing 5% methanol. A voltage of 25 kV was typically applied at the injection. The outlet of the capillary was tapered to ca. 15  $\mu\text{m}$  i.d. using a laser puller (Sutter Instruments, Novato, CA, USA). Mass spectra were acquired with dwell times of 3.0 ms per step of 1  $m/z$  unit in full-mass-scan mode. For CZE-ES-MS/MS experiments, about 30 nL sample was introduced using 300 mbar for 0.1 min. The MS/MS data were acquired with dwell 30 times of 1.0 ms per step of 1  $m/z$  unit. Fragment ions formed by collision activation of selected precursor ions with nitrogen in the RF-only quadrupole collision cell, were mass-analyzed by scanning the third quadrupole. Collision energies were typically 60 eV (laboratory frame of reference).

**Mouse Colonization**

Specific Pathogen free Female CD1 mice were purchased from Charles Rivers Laboratories, Montreal when they were 6-8 weeks old. Mice were maintained  
5 and used in accordance with the recommendations of the Canadian Council on Animal Care, Guide to the Care and Use of Experimental Animals (1993). Mice were inoculated with bacteria harvested from 36h broth culture. Aliquots of 0.2 ml, containing approximately  $10^8$  bacteria resuspended in PBS were given by gavage directly into the gastric lumen using a 20g gavage needle. Three inocula  
10 were given over a period of 6 days. No attempt was made to neutralize gastric acidity prior to inoculation. To recover viable bacteria from the stomach, mice were killed by CO<sub>2</sub> asphyxiation, and their stomachs removed whole. Stomachs were cut open along the greater curvature, and the exposed luminal surface was gently irrigated with 10 ml of sterile PBS, delivered via a syringe fitted with a 20g  
15 gavage needle, to dislodge the loosely adherent stomach contents. This step effectively diminished the small numbers of ubiquitous contaminating bacteria that otherwise overgrow on GSS agar to thereby mask the presence of the slower growing *H. pylori* organisms. The washed stomach tissue was then homogenised, and serial dilutions plated on GSS agar. *H. pylori* colonies were  
20 counted following 3-6 days incubation.

**Detection of *H. pylori* specific antibodies by ELISA**

Sera for antibody determinations were prepared from clotted blood obtained from a lateral tail vein during the course of an experiment or by cardiac puncture at the  
25 time of necropsy. Sera were screened for the presence of specific IgG isotype anti- *H. pylori* antibodies by ELISA essentially by the method of Engvall *et al* (*J. Immunol.* 109: 129-135 (1972)). Briefly, microtitre plates (Dynatech Immunolon II) were coated with 100 µl antigen (50 µg protein/ml in 0.05M carbonate buffer pH 9.8) and incubated overnight at 4°C. Antigen was prepared by resuspending  
30 plate grown *H. pylori* in PBS and sonicating the suspension until a translucent solution was obtained. The sonicate was membrane filter sterilized through a 0.45 µm filter. The protein content of the filtrate was determined by Lowry assay using a commercial kit. Sodium azide was added to 0.05% w/v and the antigen

solution was stored at 4°C. When LPS was used as the coating antigen the concentration was 10 $\mu$ g/ml. Sera were screened at a starting dilution of 1/40 and were titrated through a two-fold dilution series down a column of 8 wells. The developing antibody was goat-anti-mouse IgG conjugated to alkaline phosphatase (Caltag Laboratories). Titres were determined from plots of absorbance at 410 nm *versus* dilution and were defined as the reciprocal of the dilution giving an  $A_{410}$  equivalent to 0.25. Standard negative and positive control sera identified by a preliminary ELISA of candidate samples were included on each plate. Titres were analysed statistically by Mann Whitney Rank Sum Test and were considered to be significantly different to comparative samples when p values <0.05 were obtained.

Although various particular embodiments of the present invention have been described hereinbefore for purposes of illustration, it would be apparent to those skilled in the art that numerous variations may be made thereto without departing from the spirit and scope of the invention, as defined in the appended claims.

## SEQUENCE LISTING

<110> National Research Council of Canada  
 Logan, Susan M.  
 Wakarchuk, Warren  
 Conlan, W.  
 Monteiro, Mario A.  
 Altman, Eleonora  
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<120> Glycosyltransferases of Helicobacter pylori as a new target in prevention and treatment of H. pylori infections

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Phe Phe Asn Ser		
370		

<210> 5  
 <211> 849  
 <212> DNA  
 <213> Helicobacter pylori

<400> 5  
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 atcgccaagc ttttagaatg cgaaaaacat tttgaaatca ttccctcttt tgaaaatgtc 180  
 cctgcttttt atgaccttaa aaaacaaggc gtttttttgg cgatgaagga ttttttatgg 240  
 ttgttaaaag cgattaaaaa gcatcaaattc aaacgttga ttttggaaaaa acaggatttt 300  
 agaagcactt ttttagccaa attcattccc ataaccactc caaataaaaga aattaaaaac 360  
 gtttatccaa accgccagga gttgttttct caaattttatg ggcattgttt tgataacccc 420  
 ccatatccca tgaattttaaa aaaccccaaa aagattttga tcaacccttt cacaagatcc 480  
 atagaccgaa gttatccccctt agagcattta caaatcgttt taaaactttt aaaaccccttt 540  
 tggttacgc ttttagattt tgaagaacga tacgctttt taaaagacag agtcgcctcat 600  
 tatcgcgcta aaaccaggttt agaagaagtt aaaaacctga ttttagaaag cgatttttat 660  
 ataggagggg attcggtttt gatccatttgc gtttactatt taaagaaaaaa ttatccatc 720  
 ttttttataa gggataatga tgatccatg ccgccttataa gtaagaataa aaatttctta 780  
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 ctattataa 849

<210> 6  
 <211> 282

&lt;212&gt; PRT

&lt;213&gt; Helicobacter pylori

&lt;400&gt; 6

Met	His	Val	Ala	Cys	Leu	Leu	Ala	Leu	Gly	Asp	Asn	Leu	Ile	Thr	Leu
1				5				10					15		

Ser	Leu	Leu	Lys	Glu	Ile	Ala	Phe	Lys	Gln	Gln	Gln	Pro	Leu	Lys	Ile
					20			25					30		

Leu	Gly	Thr	Arg	Leu	Thr	Leu	Lys	Ile	Ala	Lys	Leu	Leu	Glu	Cys	Glu
					35			40					45		

Lys	His	Phe	Glu	Ile	Ile	Pro	Leu	Phe	Glu	Asn	Val	Pro	Ala	Phe	Tyr
					50			55			60				

Asp	Leu	Lys	Lys	Gln	Gly	Val	Phe	Leu	Ala	Met	Lys	Asp	Phe	Leu	Trp
					65			70		75		80			

Leu	Leu	Lys	Ala	Ile	Lys	Lys	His	Gln	Ile	Lys	Arg	Leu	Ile	Leu	Glu
					85			90			95				

Lys	Gln	Asp	Phe	Arg	Ser	Thr	Phe	Leu	Ala	Lys	Phe	Ile	Pro	Ile	Thr
					100			105			110				

Thr	Pro	Asn	Lys	Glu	Ile	Lys	Asn	Val	Tyr	Gln	Asn	Arg	Gln	Glu	Leu
					115			120			125				

Phe	Ser	Gln	Ile	Tyr	Gly	His	Val	Phe	Asp	Asn	Pro	Pro	Tyr	Pro	Met
					130			135			140				

Asn	Leu	Lys	Asn	Pro	Lys	Lys	Ile	Leu	Ile	Asn	Pro	Phe	Thr	Arg	Ser
					145			150			155		160		

Ile	Asp	Arg	Ser	Ile	Pro	Leu	Glu	His	Leu	Gln	Ile	Val	Leu	Lys	Leu
					165			170			175				

Leu	Lys	Pro	Phe	Cys	Val	Thr	Leu	Leu	Asp	Phe	Glu	Glu	Arg	Tyr	Ala
					180			185			190				

Phe	Leu	Lys	Asp	Arg	Val	Ala	His	Tyr	Arg	Ala	Lys	Thr	Ser	Leu	Glu
					195			200			205				

Glu	Val	Lys	Asn	Leu	Ile	Leu	Glu	Ser	Asp	Leu	Tyr	Ile	Gly	Gly	Asp
					210			215			220				

Ser	Phe	Leu	Ile	His	Leu	Ala	Tyr	Tyr	Leu	Lys	Lys	Asn	Tyr	Phe	Ile
					225			230			235		240		

Phe	Phe	Tyr	Arg	Asp	Asn	Asp	Asp	Phe	Met	Pro	Pro	Asn	Ser	Lys	Asn
					245			250			255				

Lys	Asn	Phe	Leu	Lys	Ala	His	Lys	Ser	His	Ser	Ile	Glu	Gln	Asp	Leu
					260			265			270				

Ala Lys Lys Phe Arg His Leu Gly Leu Leu  
275   280

<210> 7  
<211> 1050  
<212> DNA  
<213> Helicobacter pylori

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tttatcttag tggggcccaac cattacttgc gaactttca aaaaagatga aaaaatagaa 180  
gccgtttta tagacaacac caaaaaatcc ttttcaggc tgctagccat tcacaaactc 240  
gctaaaaaaaa tagggcggtt cgatatacg atcaactttaa acaaccattt ctattccgct 300  
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tgcttttat ttgcgaatt tttagaaaaaa gaattggatc aaaaaagcgt tttaccctta 480  
aaactggcct ttaaccccc cactcacacc ccaaacaccc ctaaaaaaat cggctttaac 540  
cctagcgcaa gctatggag tgctaaaaga tggccagctt cttattacgc tgaagttct 600  
gctgtttgt tagaaaaaagg gcatgaaatt tatttttttg gggctaaaga agacgctatc 660  
gtttctgaag aaattttaaa actcatcaaa ggctcattaa aaaacccctt attgtttccat 720  
aacgcttaca atctgtgcgg gaaaacaagc attgaagaat tgatagagcg catcgctgtt 780  
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aaaacgatcg tattgaacca ccatttaagc tgcgcgcctt gcaagaaaacg agtttgcct 960  
ttaaaaatcg caaaaaaaaa tttgtgcgtatc aaatctatca cgcccccttga agtccctagaa  
1020  
gcccgttcaca ctcttttaga agagccttaa  
1050

<210> 8  
<211> 349  
<212> PRT  
<213> Helicobacter pylori

<400> 8  
Met Ser Val Asn Ala Pro Lys Arg Met Arg Ile Leu Leu Arg Leu Pro  
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Asn Trp Leu Gly Asp Gly Val Met Ala Ser Ser Leu Phe Tyr Thr Leu  
20   25   30  
  
Lys His His Tyr Pro Asn Ala His Phe Ile Leu Val Gly Pro Thr Ile  
35   40   45  
  
Thr Cys Glu Leu Phe Lys Lys Asp Glu Lys Ile Glu Ala Val Phe Ile  
50   55   60  
  
Asp Asn Thr Lys Lys Ser Phe Phe Arg Leu Leu Ala Ile His Lys Leu  
65   70   75   80  
  
Ala Gln Lys Ile Gly Arg Cys Asp Ile Ala Ile Thr Leu Asn Asn His  
85   90   95  
  
Phe Tyr Ser Ala Phe Leu Leu Tyr Ala Thr Lys Thr Pro Val Arg Ile  
100   105   110

Gly Phe Ala Gln Phe Phe Arg Ser Leu Phe Leu Ser His Ala Ile Ala  
 115 120 125

Pro Ala Pro Lys Glu Tyr His Gln Val Glu Lys Tyr Cys Phe Leu Phe  
 130 135 140

Ser Gln Phe Leu Glu Lys Glu Leu Asp Gln Lys Ser Val Leu Pro Leu  
 145 150 155 160

Lys Leu Ala Phe Asn Leu Pro Thr His Thr Pro Asn Thr Pro Lys Lys  
 165 170 175

Ile Gly Phe Asn Pro Ser Ala Ser Tyr Gly Ser Ala Lys Arg Trp Pro  
 180 185 190

Ala Ser Tyr Tyr Ala Glu Val Ser Ala Val Leu Leu Glu Lys Gly His  
 195 200 205

Glu Ile Tyr Phe Phe Gly Ala Lys Glu Asp Ala Ile Val Ser Glu Glu  
 210 215 220

Ile Leu Lys Leu Ile Lys Gly Ser Leu Lys Asn Pro Ser Leu Phe His  
 225 230 235 240

Asn Ala Tyr Asn Leu Cys Gly Lys Thr Ser Ile Glu Glu Leu Ile Glu  
 245 250 255

Arg Ile Ala Val Leu Asp Leu Phe Ile Thr Asn Asp Ser Gly Pro Met  
 260 265 270

His Val Ala Ala Ser Met Gln Thr Pro Leu Ile Ala Leu Phe Gly Pro  
 275 280 285

Thr Asp Glu Lys Glu Thr Arg Pro Tyr Lys Ala Gln Lys Thr Ile Val  
 290 295 300

Leu Asn His His Leu Ser Cys Ala Pro Cys Lys Lys Arg Val Cys Pro  
 305 310 315 320

Leu Lys Asn Ala Lys Asn His Leu Cys Met Lys Ser Ile Thr Pro Leu  
 325 330 335

Glu Val Leu Glu Ala Ala His Thr Leu Leu Glu Glu Pro  
 340 345

<210> 9  
<211> 822  
<212> DNA  
<213> Helicobacter pylori

<400> 9  
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gatgcgattt attctaaaac ttttgaaggc gggttgcacc ccttagtgaa aaagcattta 180  
cacccttatt tcattcacgca aaacatcaa gacatgggaa ttacaaccag tctcatcagt 240  
gaggTTTcta agttttatta cgctttaaaa taccatgcga agtttatgag cttggggagag 300  
cttgggtgct atgcgagcca ttattcccttg tgggaaaaat gcatagaact caatgaagcg 360  
atctgtatTT tagaagacga tataaccttg aaagaggatt ttaaagaggg cttggatTTT 420  
ttagaaaaac acatccaaga gttaggctat gttcgcttga tgcatttatt atatgatccc 480  
aatattaaaa gtgagccatt gaaccataaa aaccacgaga tacaagagcg tgttagggatt 540

attaaagctt atagcgaagg ggtggggact caaggctatg tgatcacgcc caagattgcc 600  
 aaagtttta aaaaacacag cggaaaatgg gttgttcctg tggatacgat aatggacgct 660  
 actttatcc atggcgtgaa aaatctggg ttacaacctt ttgtgatcgc tgatgatgag 720  
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 gaactccatt taaaatattt gaaatatgg cagtttatat ag 822

<210> 10  
 <211> 273  
 <212> PRT  
 <213> Helicobacter pylori

<400> 10
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Thr His His Gln Val Gln Ile Phe Asp Ala Ile Tyr Ser Lys Thr Phe
35 40 45
Glu Gly Gly Leu His Pro Leu Val Lys Lys His Leu His Pro Tyr Phe
50 55 60
Ile Thr Gln Asn Ile Lys Asp Met Gly Ile Thr Thr Ser Leu Ile Ser
65 70 75 80
Glu Val Ser Lys Phe Tyr Tyr Ala Leu Lys Tyr His Ala Lys Phe Met
85 90 95
Ser Leu Gly Glu Leu Gly Cys Tyr Ala Ser His Tyr Ser Leu Trp Glu
100 105 110
Lys Cys Ile Glu Leu Asn Glu Ala Ile Cys Ile Leu Glu Asp Asp Ile
115 120 125
Thr Leu Lys Glu Asp Phe Lys Glu Gly Leu Asp Phe Leu Glu Lys His
130 135 140
Ile Gln Glu Leu Gly Tyr Val Arg Leu Met His Leu Leu Tyr Asp Pro
145 150 155 160
Asn Ile Lys Ser Glu Pro Leu Asn His Lys Asn His Glu Ile Gln Glu
165 170 175
Arg Val Gly Ile Ile Lys Ala Tyr Ser Glu Gly Val Gly Thr Gln Gly
180 185 190
Tyr Val Ile Thr Pro Lys Ile Ala Lys Val Phe Lys Lys His Ser Arg
195 200 205
Lys Trp Val Val Pro Val Asp Thr Ile Met Asp Ala Thr Phe Ile His
210 215 220
Gly Val Lys Asn Leu Val Leu Gln Pro Phe Val Ile Ala Asp Asp Glu
225 230 235 240

Gln Ile Ser Thr Ile Ala Arg Lys Glu Gln Pro Tyr Ser Pro Lys Ile  
 245 250 255

Ala Leu Met Arg Glu Leu His Phe Lys Tyr Leu Lys Tyr Trp Gln Phe  
 260 265 270

Ile

<210> 11  
 <211> 1120  
 <212> DNA  
 <213> Helicobacter pylori

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 agtgaaaaac tttttataa aatccactgc ctggtagata acttaaggct tgaaaaccag 180  
 tgcaaattga aagaaactct agcccccttta agcgcttta tgagcgtgga ttttttagac 240  
 attcaaccc ctaatctta caccccttca atagaacccct ctgcgattga taaaatcaat 300  
 gaagctttt tgcaactcaa ttttacgct aagactcgct ttctaaaat ggtcatgtgc 360  
 cgcttgtttt tggcttctt atccccgcaa tacgacaaaa tcatcatgtt tgatgcggac 420  
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 1120

<210> 12  
 <211> 372  
 <212> PRT  
 <213> Helicobacter pylori

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 35 40 45

His Cys Leu Val Asp Asn Leu Ser Leu Glu Asn Gln Cys Lys Leu Lys  
 50 55 60

Glu Thr Leu Ala Pro Phe Ser Ala Phe Met Ser Val Asp Phe Leu Asp  
 65 70 75 80

Ile Ser Thr Pro Asn Leu Tyr Thr Pro Ser Ile Glu Pro Ser Ala Ile  
 85 90 95  
 Asp Lys Ile Asn Glu Ala Phe Leu Gln Leu Asn Ile Tyr Ala Lys Thr  
 100 105 110  
 Arg Phe Ser Lys Met Val Met Cys Arg Leu Phe Leu Ala Ser Leu Phe  
 115 120 125  
 Pro Gln Tyr Asp Lys Ile Ile Met Phe Asp Ala Asp Thr Leu Phe Leu  
 130 135 140  
 Asn Asp Val Ser Glu Ser Phe Phe Ile Pro Leu Asp Gly Tyr Tyr Phe  
 145 150 155 160  
 Gly Ala Ala Lys Asp Phe Ser Ser Pro Lys Asn Leu Lys His Phe Gln  
 165 170 175  
 Thr Glu Arg Glu Arg Glu Pro Arg Gln Lys Phe Phe Leu His Glu His  
 180 185 190  
 Tyr Leu Lys Glu Lys Asp Met Lys Ile Ile Cys Glu Asn His Tyr Asn  
 195 200 205  
 Val Gly Phe Leu Ile Val Asn Leu Lys Leu Trp Arg Ala Asp His Leu  
 210 215 220  
 Glu Glu Arg Leu Leu Asn Leu Thr His Gln Lys Gly Gln Cys Val Phe  
 225 230 235 240  
 Cys Pro Glu Gln Asp Ile Leu Thr Leu Ala Cys Tyr Gln Lys Val Leu  
 245 250 255  
 Gln Leu Pro Tyr Ile Tyr Asn Thr His Pro Phe Met Val Asn Gln Lys  
 260 265 270  
 Arg Phe Ile Pro Asn Lys Lys Glu Ile Val Met Leu His Phe Tyr Phe  
 275 280 285  
 Val Gly Lys Pro Trp Val Leu Pro Thr Ala Leu Tyr Ser Lys Glu Trp  
 290 295 300  
 His Glu Thr Leu Leu Lys Thr Pro Phe Tyr Ala Glu Tyr Ser Val Lys  
 305 310 315 320  
 Phe Leu Lys Gln Met Thr Glu Phe Leu Ser Leu Lys Asp Lys Gln Lys  
 325 330 335  
 Thr Phe Glu Phe Leu Ala Pro Leu Leu Asn Lys Lys Thr Leu Leu Glu  
 340 345 350  
 Tyr Val Phe Phe Arg Leu Asn Arg Ile Phe Lys Arg Leu Lys Glu Lys  
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 Leu Leu Asn Ser  
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<210> 13  
<211> 843  
<212> DNA

<213> Helicobacter pylori

<400> 13

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cccgcttttt atgacccat tttttttttgg cgatgaagga ttttttatgg 240
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<210> 14

<211> 280

<212> PRT

<213> Helicobacter pylori

<400> 14

Met His Val Ala Cys Leu Leu Ala Leu Gly Asp Asn Leu Ile Thr Leu			
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20	25	30	

Leu Gly Thr Arg Leu Thr Leu Lys Ile Ala Lys Leu Leu Glu Cys Glu			
35	40	45	

Lys His Phe Glu Ile Ile Pro Val Phe Lys Asn Ile Pro Ala Phe Tyr			
50	55	60	

Asp Leu Lys Lys Gln Gly Val Phe Trp Ala Met Lys Asp Phe Leu Trp			
65	70	75	80

Leu Leu Lys Ala Leu Lys Lys His Lys Ile Lys His Leu Ile Leu Glu			
85	90	95	

Lys Gln Asp Phe Arg Ser Ala Leu Leu Ser Lys Phe Val Ser Ile Thr			
100	105	110	

Thr Pro Asn Lys Glu Ile Lys Asn Ala Tyr Gln Asn Arg Gln Glu Leu			
115	120	125	

Phe Ser Gln Ile Tyr Gly His Val Phe Asp Asn Ser Gln Tyr Ser Met			
130	135	140	

Ser Leu Lys Asn Pro Lys Lys Ile Leu Ile Asn Pro Phe Thr Arg Glu			
145	150	155	160

Asn Asn Arg Asn Ile Ser Leu Glu His Leu Gln Ile Val Leu Lys Leu			
165	170	175	

Leu Lys Pro Phe Cys Val Thr Leu Leu Asp Phe Glu Glu Arg Tyr Ala			
180	185	190	

Phe Leu Lys Asp Glu Val Ala His Tyr Arg Ala Lys Thr Ser Leu Glu  
 195 200 205

Glu Ala Lys Asn Leu Ile Leu Glu Ser Asp Leu Tyr Ile Gly Gly Asp  
 210 215 220

Ser Phe Leu Ile His Leu Ala Tyr Tyr Leu Lys Lys Asn Tyr Phe Ile  
 225 230 235 240

Phe Phe Tyr Arg Asp Asn Asp Asp Phe Met Pro Pro Lys Asn Glu Asn  
 245 250 255

Phe Leu Lys Ala His Lys Ser His Phe Ile Glu Gln Asp Leu Ala Thr  
 260 265 270

Gln Phe Arg His Leu Gly Leu Leu  
 275 280

<210> 15

<211> 850

<212> DNA

<213> Helicobacter pylori

<400> 15

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 atcgccaaagc ttttagaatg cgaaaaacat tttgaatca ttccatttt tgaaaatatc 180  
 cctgctttttt atgatcttaa aaaacaaggc gttttttggg cgatgaagga ttttttatgg 240  
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 ccatacttcca tgaattttaaa aaacccctaa aagatttga tcaaccccttt cacaagatcc 480  
 atagagcgaa gtatccccctt agagcattta aaaatcgctt taaaactctt aaaaccccttt 540  
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 tatacggtcta aaaccagttt agaagaagtt aaaagcctga ttttagaaaa cgatttttat 660  
 ataggggggg attcgttttt aatccatttg gtttactatt taaaagaaaaa ttattttatac 720  
 ttttttata gggataatga cgatttcatg ccacctaattt gtaagaagga aaattttcta 780  
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 cttattataa 850

<210> 16

<211> 283

<212> PRT

<213> Helicobacter pylori

<400> 16

Met His Val Ala Cys Leu Leu Ala Leu Gly Asp Asn Leu Ile Thr Leu  
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 20 25 30

Leu Gly Thr Arg Leu Thr Leu Lys Ile Ala Lys Leu Leu Glu Cys Glu  
 35 40 45

Lys His Phe Glu Ile Ile Pro Ile Phe Glu Asn Ile Pro Ala Phe Tyr  
 50 55 60

Asp Leu Lys Lys Gln Gly Val Phe Trp Ala Met Lys Asp Phe Leu Trp  
65 70 75 80

Leu Leu Lys Ala Ile Lys Lys His Lys Ile Lys His Leu Ile Leu Glu  
85 90 95

Lys Gln Asp Phe Arg Ser Phe Leu Leu Ser Lys Phe Val Ser Ile Thr  
100 105 110

Thr Pro Asn Lys Glu Ile Lys Asn Val Tyr Gln Asn Arg Gln Glu Leu  
115 120 125

Phe Ser Pro Ile Tyr Gly His Val Phe Asp Asn Pro Pro Tyr Pro Met  
130 135 140

Asn Leu Lys Asn Pro Lys Lys Ile Leu Ile Asn Pro Phe Thr Arg Ser  
145 150 155 160

Ile Glu Arg Ser Ile Pro Leu Glu His Leu Lys Ile Val Leu Lys Leu  
165 170 175

Leu Lys Pro Phe Cys Val Thr Leu Leu Asp Phe Glu Glu Arg Tyr Ala  
180 185 190

Phe Leu Gln Asn Glu Ala Thr His Tyr Arg Ala Lys Thr Ser Leu Glu  
195 200 205

Glu Val Lys Ser Leu Ile Leu Glu Ser Asp Leu Tyr Ile Gly Gly Asp  
210 215 220

Ser Phe Leu Ile His Leu Ala Tyr Tyr Leu Lys Lys Asn Tyr Phe Ile  
225 230 235 240

Phe Phe Tyr Arg Asp Asn Asp Asp Phe Met Pro Pro Asn Gly Lys Lys  
245 250 255

Glu Asn Phe Leu Lys Ala His Lys Ser His Tyr Ile Glu Gln Asp Leu  
260 265 270

Ala Lys Lys Phe Arg His Leu Gly Leu Ile Ile  
275 280